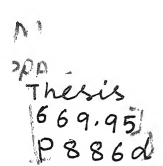
# DEVELOPMENT OF MAGNETIC ALLOYS BY THE DIRECT REDUCTION OF OXIDES

By
SHARDA NAND PRASAD

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1005





DEPARTMENT OF METALLURGICAL ENGINEERING

INDIAN INSTITUTE OF TECHNOLOGY KANPUR APRIL, 1986

# DEVELOPMENT OF MAGNETIC ALLOYS BY THE DIRECT REDUCTION OF OXIDES

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in Partial Fulfilment of the Requirements
for the Degree of
MASTER OF TECHNOLOGY

By
SHARDA NAND PRASAD

to the

DEPARTMENT OF METALLURGICAL ENGINEERING
INDIAN INSTITUTE OF TECHNOLOGY KANPUR
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#### CERTIFICATE

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REDUCTION OF OXIDES', has been carried out by
Mr. Sharda Nand Prasad under my supervision and
it has not been submitted elsewhere for the
award of a degree.

April 1936.

K.N. Rai

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April, 1936

SHARDA NAND PRASAD

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#### ABSTRACT

The present investigation involves the study of process variables on the development of magnetic properties in Alnico alloys. These process variables are fabrication method of alloys, temperature of annealing, time of annealing, time and temperature of magnetic ageing, and addition of alloying elements (Cr. Co and Cu).

Alloys chosen for study contain 70-57% Fe, 20-28% Ni, 5% Al and 5-10% Cr in Fe-Ni-Al-Cr system and 63-52% Fe, 20-12% Ni, 10% Al, 5-20% Co and 2-6% Cu in Fe-Ni-Al-Co-Cu system. All alloys were annealed at 1000°C for 3 hrs. and then tempered at 650°C for  $3\frac{1}{2}$  hrs. in H<sub>2</sub> atmosphere.

The microstructural features of Alnico alloys were studied by optical microscope. The phase analysis was carried out by using X-ray diffraction technique.

Microhardness and Resisvity both were found to increase linearly with increased amount of alloying elements.

Addition of Co and Cu was aimed at increasing  $M_s$ ,  $M_r$  and iHc values. Chromium increases iHc, but lowers  $M_s$  and  $M_r$  values,  $H_2$  annealing appears to improve  $M_s$ ,  $M_r$  and iHc values.

Magnetic ageing of alloys in the field of 1200 Gauss for 1/2 hr. at 650°C showed promising improvement in  $\rm ^M_S$  ,  $\rm ^M_r$  and  $\rm ^iHc$  values and induced loop squareness.

#### CHAPTER - I

#### INTRODUCTION AND LITERATURE REVIEW

#### ABSTRACT:

This chapter reviews the subject matters related to structure, properties, the general methods of fabrication, characterization and applications of Alnico magnet alloys.

#### 1.1 Introduction:

of industrial devices and development of modern instruments necessitated the development of various kinds of magnetic materials. The relation among magnetic properties composition, crystal structure and the method of fabrication, has immense important. As a consequence this has stimulated much scientific and technological development work aimed at influencing the magnetic properties in order to develop materials with most advantageous combination for applications in telecommunication and electrical engineering.

The discoveries in this field are Alnico allows as permanent magnet by Mishima in 1931,

ferromagnetic resonance by Griffith in 1946, antiferromagnetism by Neel in 1948, theory of single domain particles by Wolliams and Bozorth in 1949, magnetic ferrite by Snock in 1949 and so on.

A permanent magnet is a storage device. The requirement of every saving as well as miniaturisation of devices employing magnets has generated in recent years a renewed interest in the development of new, high energy product magnetic materials. Wide spread interest in Alnico alloys which had anisotropy constant  $(K_1)$  in the range of  $(.3 \times 10^6 - 1.9 \times 10^6) \, \mathrm{ergs/cm}^3$  which is much larger than that of steels. Properties of some important permanent magnets are in Table (1.1).

A high quality permanent magnet material must have the following properties: (i) high saturation magnetization ( $M_r$ ), (ii) high curie temperature ( $T_c$ ), (iii) Uni-axial anisotropy (iv) larger value of magneto crystalline anisotropy constant ( $K_1$ ), (v) high coercive force ( $H_c$ ), (vi) a large fulness factor and (vii) low permanent permeability.

Besides these requirements, the material must have high corrossion resistant, thermal stability and should be cheap.

The main contribution towards the saturation  $magnetization (M_s)$  of alnico magnet alloys is made by

TABLE : 1.1

Properties of	some Permanent	Magnet Ma	terials	
Materials	Composition (weight %)	Rema- nence B r (Gauss)	Coer- civity <sup>H</sup> c Orsted	Maximum energy product (BH) max M.G.Oe
36 Co Steel	36 Co, 3.75 w, 5.75 Cr, 0.8 C	9600	228	.93
Alnico 1	12 Al, 23 Ni, 5 Co, 60 Fe	6600	<b>54</b> 0	1.4
Alnico 2	10 Al, 18 Ni, 13 Co, 6 Cu, 63 Fe	7000	6 50	1.7
Alnico 3	12 Al, 26 Ni, 3 Cu, 59 Fe	· 6 <b>40</b> 0	560	1.4
Alnico 4	12 Al, 28 Ni, 5 Co, 55 Fe	5500	<b>73</b> 0	1.4
Alnico 5#	8 Al, 15 Ni, 24 Co, 3 Cu, 50 Fe	12000	<b>72</b> 0	5 <b>.</b> 0
Alnico 5	8 Al, 15 Ni, 24 Co, 3 Cu, 50 Fe	13100	700	6.5
Alnico 6#	8 Al, 17 Ni, 23 Co, 3 Cu, 45 Fe, 4 Ti	7 500	975	2.8
Alnico 8#	8 Al, 14 Ni, 38 Co, 3 Cu, 29 Fe, 8 Ti	7100	2000	5.5
Alnico 9 <sup>#</sup>	7 Al, 15 Ni, 35 Co, 4 Cu, 34 Fe, 5 Ti	10400	1600	8.5

<sup>\*</sup> Anisotropic properties

parallel (i.e. ferromagnetically coupled) magnetic moment of Fe atoms in the ferromagnetic phase  $\alpha$  (Fe38 Ni Al) and in weakly magnetic phase  $\alpha^*$  (Fe6 Ni8 Al7). Both phases have Body-centred crystal structure (1). The alnico alloys have body-centred crystal structure with unixial anisotropy i.e. due to easy direction of magnetization is in the [100] direction as shown in Fig. (1.1).

The coercive force of alnico alloys depends on many factors. High coercive force is due to very large anisotropy fields varying with composition of alloys. The coercive force finally developed in a magnet is however, influenced by many processing perameters (2). High coercive force can be developed by making conditions favourable for pinning domain walls. This approach has been used in some alnico alloys. Yet another method to increase coercivity is by precipitating out elongated particles of the magnetic phase in a non-magnetic matrix, thereby coupling high magnetocrystalline anisotropy field as well as shape anisotropy effect to yield high value of coercivity.

# 1.2 Alnico Alloys:

The alloys of the Fe-Ni-Al system with or without specific alloying additions, are known as Alnico alloys. Depending upon composition, history of

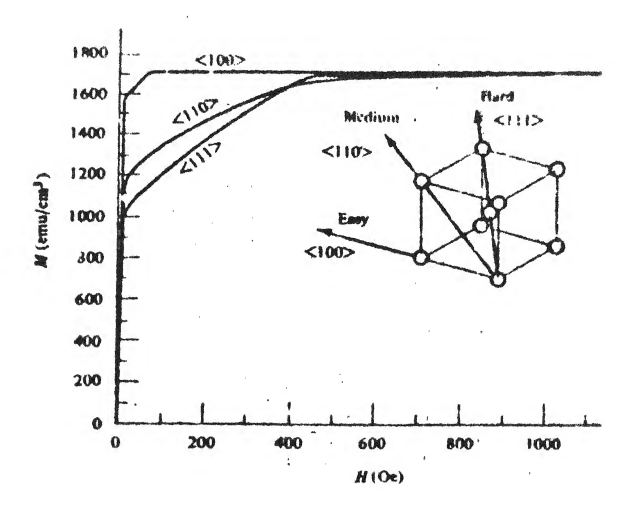


Fig. 1.1: Magnetization curves for single of the from

of fabrication and purpose of development, they may be further classified as Alnico - 1, 2, 3, 4, 5, 5 DG, 6, 7, 8 and 9 etc. (3). They have wide range of magnetic properties and are suitable for using as permanent magnets. These alloys are brittle and must be cast to shape. Details of the optimum composition and heat treatment of these isotropic alloys in which magnetic properties are independent of orientation, have been studied by Betteridge (4). The best properties were obtained for an alloy containing 60% Fe - 28% Ni - 12% Al which had been quenched at 28°C per second from the single phase state at 1100°C and tempered for 4 hours at 650°C. This treatment gave a coercivity (BH\_c)=515 Orsted and a maximum energy product  $(BH)_{max} = 1.35 \times 10^6$ G. Oe. The coercivity was shown to depend very critically on the Al content while the remanence depended more on the Ni content.

The magnetic properties of Alnico alloys can be improved by the addition of Co, Cu and other alloying elements. Betteridge (4) also investigated the effect of addition Cu to (Fe - Ni - Al) system alloys and found that a 3.5 weight percent Cu addition increased (BH)<sub>max</sub> to 1.5 x 10<sup>6</sup> G.Oe after quenching from 1100°C and tempered at 550°C. The Cu addition increased the rate of precipitation of phases so that alloys required more rapid cooling.

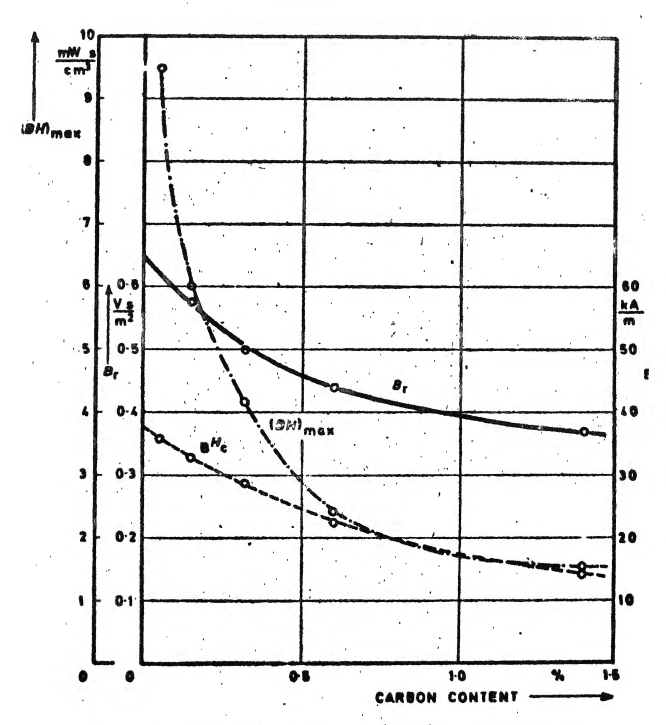


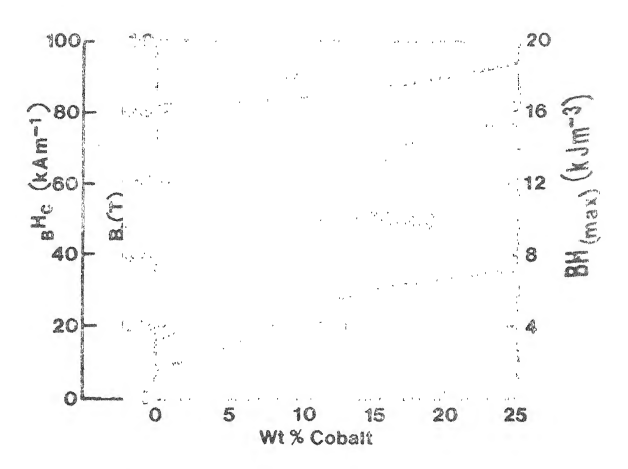
Fig. 1.2: Effect of carbon on the magnetic properties of Fe-Ni-Al - system alloy.

The effect of elastic stress on the precipitation of phases and addition of Cu and Ti on magnetic properties of Fe - Ni - Al system alloys have been investigated by Yermolenko and Korolyov (5) who obtained improved optimum permanent magnetic properties.

The carbon content of Fe - Ni - Al system alloys is kept as low as possible, since M<sub>2</sub> and intrinsic coercivity (<sup>i</sup>H<sub>C</sub>) are seriously reduced with carbon content even below .1%. However, small additions of Ti and Cr (about 0.5%) are made to combine with carbon and counteract its deleterious effect. The effect of carbon on magnetic properties of Fe - Ni - Al system alloys is shown in Fig. (1.2), observed by Betteridge (4).

Betteridge and Zumbush (6) had found that the magnetic properties of Alnico alloys could be improved by the addition of Co as shown in Fig. (1.3). The gains were progressive as the Co content was increased to as high as 20 - 25 %, although the benefits may become uneconomic at higher Co content. A higher solution temperature of about (1250°C - 1300°C) is required. The critical cooling rate varies with composition. The ageing temperature remains in the range of 600°C - 650°C.

The composition and magnetic properties of some known Alnico magnet alloys are given in Table (1.2).



Pir. 1.3: Dependence of ...., remandage and carry product on achilt content of Fe-Ni-Al rystem alloy

TABLF: 1.2

Compositions and magnetic properties of isotropic

Alnico magnets.

Com	posi	tion	wt	: ,3")	Remanence (B_) Gauss	Coercive force ( <sup>B</sup> H <sub>C</sub> )	(BH) max G.Oe x 10 <sup>3</sup>
(Ba	lanc	e Fe	.)			Orsted	
Ni	Co	Al	Cu	Ti			
24	0	12	2	0	5000	502	12 57
21	3	11	3	1	5500	530	1383
16	12	9	4	1	6 500	620	1635
18	17	8	4	4	<b>7</b> 000	725	1760

### 1.3 Anisotropic Alnico Magnet Alloys:

properties of alnico alloys by imposing a strong magnetic field during cooling from the solution treatment temperature. The critical temperature range over which the magnetic field must be maintained is 900°C - 600°C. The resulting magnetic properties are anisotropic, being higher in the direction of the applied field than perpendicular to it. The improvement in properties are mainly shown by increase remanence (B<sub>r</sub>), which may become as high as .9 times the saturation flux in stead of only .6 times to isotropic alloys. This results in

a much square demagnetizing curve and a high value of  $(BH)_{max}$ .

The compositions and magnetic properties of anisotropic alnico magnet alloys are given in Table (1.3).

TABLE: 1.3

Compositions and magnetic properties of Anisotropic

Alnico magnet alloys

Composition (Wt %) (Balance Fe)					Remanence (B <sub>r</sub> ) Gauss	Coercive force (H <sub>C</sub> ) Orsted	(BH) <sub>max</sub> G.0e x 10 <sup>6</sup>
Ni	œ	Al	Cu	Ti			
12	23	7.8	2	0.5	12,000	580	5.03
14	32	7	4	4	8,000	1380	5 <b>. 2</b> 8

### 1.4 Sintered Alnico Magnet Alloys:

Planchard and Bronner (8) have produced small magnets in large quantities by powder techniques. The powder magnets have lower magnetic properties than those of cast alloys of the same basic composition, because of the presence of porosity or non-magnetic blinders. The sintered magnets can be heat treated in magnetic field to give improved anisotropic properties. The best magnetic properties are obtained when the

particle elongation occurs along the < 100 > axis lying most nearly parallel to the field direction. Typical composition and magnetic properties for the same common alnico alloys, permitting a comparison between sintered and cast materials, are given in Table (1.4).

TABLE: 1.4

Magnetic properties of cast and sintered Alnico Magnet alloys.

	.oy Co ılance		ition (W	t %)	Desig- Br C (BH) nation Gauss Orsted x 10 G.Oe	max 6
A1 10	Ni 19	Ca 12`	Cu 6		Alnico Gast 7400 528 1.59 Sintered 7000 503° 1.53	
10	21	12	9		Alnico Cast 6800 610 1.59 Sintered 6100 591 1.53	
8	11.5	21	4		Alcomax-II Cast 1270 578 4.27 Sintered 1120 566 3.27	
9	21	20	2		Hycomax Cast 8500 792 2.72 Sintered 7900 780 2.61	

# 1.5 Columnar Grained Alnico Magnet Alloys:

Wright (9) have improved the magnetic properties of the alnico magnets by explotting the difference in properties in different crystallographic directions with

the help of producing columnar grained structure in the alloys. By careful control of thermal gradient during solidification, it is possible to grow long columnar grains with the <100> axes of the alloys lying along the direction of growth. When these alloys are heat treated in a magnetic field, the magnetic properties are better than those of field treated random grain alloys.

The compositions and magnetic properties of columnar grained alnico magnets are given in Table (1.5).

TABLE: 1.5

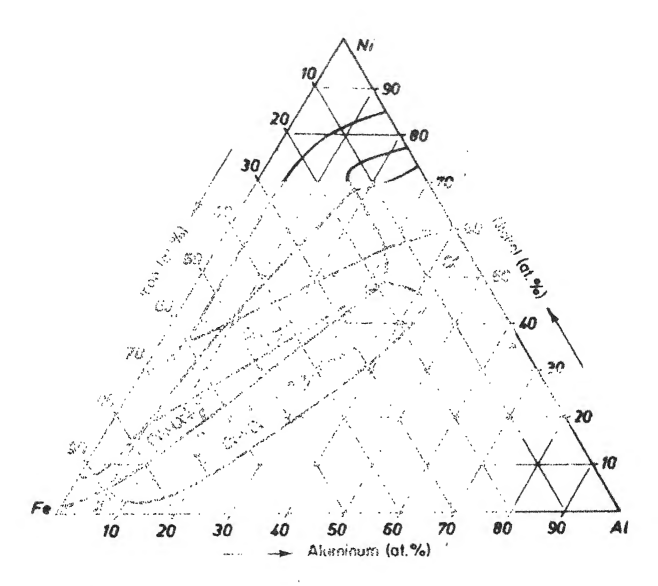
Compositions and magnetic properties of columnar

Alnico magnet alloys

Composition (Wt %) (Balance Fe)						Remanence (B <sub>r</sub> ) Gauss	Coercive force (H <sub>C</sub> ) Orsted	(BH) <sub>max</sub> G. Oe x 10
Ni	œ	Al	Cu	Мþ	Ti			-
13	24	7.8	2	1	4	13,000	704	7.04
14	32	7	4	1	4	10,000	1380	7.54

# 1.6 Phase Relation in Fe-Ni-Al System Alloys:

De Vos (10, 11) had investigated the phases present in Fe-Ni-Al system alloys using X-ray technique as shown in Fig. (1.4). One single Body-centred crystal



Pic. 1.4 : Phose dinar at fire the Fe-DE-Al'system.

structure  $\alpha$  - phase having a super-structure of the Cs Cl type exists above 1200°C. Between 1200 - 850°C, a B.C.C. $\alpha'$  (Fe<sub>38</sub> Ni Al) and another F.C.C. $\gamma_1$  phase exist.  $\gamma_1$  phase can not be supercooled to room temperature, but spontaneously transform into a B.C.C. structure known as  $\alpha$  phase (Fe<sub>6</sub> Ni<sub>8</sub> Al<sub>7</sub>) during cooling. Below 850°C, the alloy consists of two B.C.C. phases. These are  $\alpha'$  (Fe<sub>38</sub> Ni Al), rich in Fe content (ferromagnetic) and  $\alpha$  (Fe<sub>6</sub> Ni<sub>8</sub> Al<sub>7</sub>), low in Fe content (paramagnetic). After tempering for about two months at 600°C, gives rise to another F.C.C. phase ( $\gamma$ ).

# 1.7 The Effect of Heat Treatment on Magnetic Properties of Alnico Magnet Alloys:

The alnico magnet alloys show high coercivity, produced by precipitation hardening method. The stress anisotropy is introduced by high cooling rate during casting in order to obtain high coercivity. The alloy can be heat treated by the following routine: (i) quenching from high temperature (1000 - 1100°C) to room temperature and a subsequent ageing to a medium temperature (550 - 750°C). With the duration of ageing, the coercivity first increases, reaches a maximum and then falls.

The effect of temperature and time of ageing is shown in Figs. (1.5 to 1.6), as reported by Kronenberg

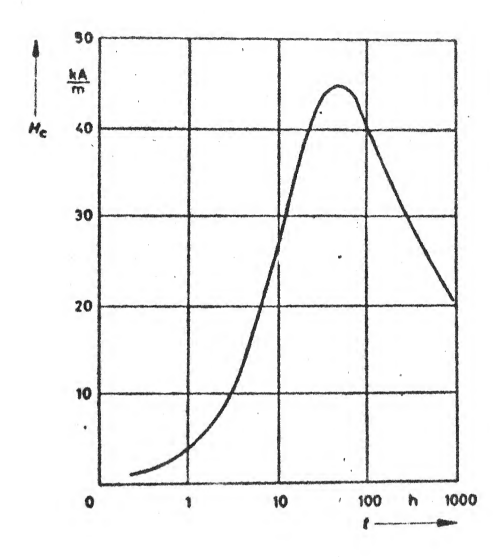


Fig. 1.5: Dependence of coercivity on duration of temperature (600°C) For Almic: 5

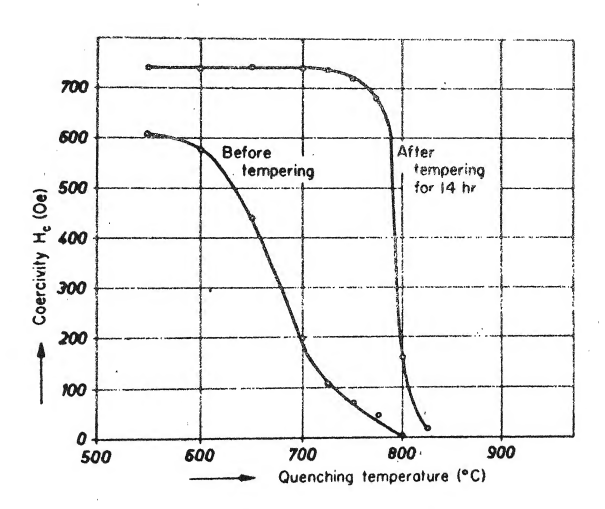


Fig. 1.6: Variation of coercivity of Alaico 5 before and after tempering as a function of quenching temperature.

and Bohlmann (12).

During ageing a fine precipitation of the insoluble particles (non-magnetic) results. The disporsion of these insoluble particles promotes optimum coercivity as a result of their pinning action on domain walls. The particles grow in size with further ageing and larger is-lands and inclusions are formed. These induce decrease in coercivity from maximum value.

#### 1.8 Effect of Magnetic Annealing:

In order to develop the magnetic properties with commercial viability, further attempts have been made to design new treatments. There are two other techniques worth mentioning for this purpose, namely the magnetic annealing and magnetic cooling both require heating or cooling essentially in a fixed magnetic field in order to facilitate alignment of magnetic domains around the curie temperature during para to ferromagnetic transformation and induce magnetic anisotropy. coercivity of the alloy depends on the degree of shape anisotropy and effective particle size. This is achieved by the spionodally decomposed microstructure, comprising of interconnected  $\alpha$  phase (paramagnetic) in a  $\alpha'$  (Fe rich) ferromagnetic matric. The application of magnetic field induces anisotropy in the bulk magnetic parameters as confirmed by Moesbaur spectroscopy (13,14).

Okade (15) has noticed the elongation of the microstructure in the direction of applied field and improvement in magnetic properties in Fe - 28% Cr - 15% Co - 1% Nb alloy.

Cremer and Pfeffier (16) also found a readily obserable induced shape anisotropy in the microstructure of an Fe - 29% Cr - 23.5% Co alloy and noticed that Mo addition enhanced the effect. The particle alignment was parallel to the <100> direction e.g. the direction of applied field.

Kaneko (17) found that particle orientation in a single crystal specimen of Fe - 30% Cr - 23% Co - 1% Si alloy, did not show preferrence for precipitation along defined crystallographic directions, although magnetic properties were marginally better for specimens, field treated in <100> direction.

The effectiveness of the magnetic annealing depends upon various parameters such as magnetic field intensity, temperature and time of annealing.

# 1.9 Effect of Annealing Atmosphere:

The annealing atmosphere is advisable to be carried out in inert or reducing atmosphere (18). The annealing atmosphere has profound effect on the magnetic properties of Fe - Ni - Al system alloys. It suggests (19) that the atmosphere probably affects the concentration

and distribution of impurity atoms through annealing and through this alters the orientation dependence of grain boundary mobility and thus influence the formation of nucleii. Another group of workers (20) atribute the due effect to an obsorbed otygen layer on the specimen surface, which causes decrease in magnetic properties.

However, many workers feel (19) that  $\rm H_2$  atmosphere annealing develops the best magnetic properties in these alloys, because  $\rm H_2$  being a strong reducing agent, may be instrumental in removing  $\rm O_2$ , C,  $\rm N_2$  or S from the matrix.

#### 1.10 Effect of Cooling Rate on Magnetic Properties:

The rate of cooling affects the grain size and coercivity of alloys. Higher rate of cooling develops stress anisotropy in the solidified alloys after casting which increases the corrcivity and remanence. The stress anisotropy constant  $(K_S)$  is defined as  $K_S = \frac{3}{2} \lambda_{Si} \times \sigma$  where  $\lambda_{Si}$  is saturation magnetostriction (assumed to be isotropic) and  $\sigma$  is uni-axial stress. For stressed particles of the solidified alloys, the coercivity  $(H_C)$  is proportional to  $\frac{\lambda_{Si} \sqrt{n}}{M_S}$  where  $M_S$  is saturation magnetization and n is the dislocation density of the solidified alloy (21).

## 1.11 The Effect of Grain Size and Inclusion on Magnetic Properties:

Besides dissolved impurities, inclusion and grain size have marked effect on magnetic properties.

Uniformity among grain size and distribution is also important as large and small grains scattering is supposed to cause greater magnetic loss because of difference in hysteris characteristics (22).

Pfeiffier (23) summarises the effect of grain size and inclusion on coercivity of Fe - Ni - Al system alloy by the relation

$$H_{C} = \frac{15}{16} \pi \frac{\gamma_{W}}{\mu_{O} M_{S}} \cdot \frac{1}{d_{G}}$$

where  $\gamma_{_{
m W}}$  is wall energy, M $_{_{
m S}}$  is saturation magnetization,  $\mu_{_{
m O}}$  is initial permeability, H $_{_{
m C}}$  is coercivity of alloy with inclusion and d $_{_{
m G}}$  is grain diameter.

Thus the coercivity of alloy with inclusion is larger than that of pure alloy of same composition.

The effect of inclusion size and distribution in Fe - Ni - Al system alloy with alloying elements such as Co, Cu and Cr etc. notes that submicron dispersion of inclusion is advisable for achieving high coercivity by heat treatment (ageing).

#### 1.12 General Methods of Fabrication:

Generally alnice magnet alloys were fabricated by induction melting technique. The earliest method used for fabrication was by solidification of liquid alloy in permanent mold during thermite process, as result of aluminothermic reaction where during solidification rod like precipitates form parallel to the cooling direction. However, the casting approach run into insurmountable difficulties and is ruled out now.

The most commonly used technique for producing alnico magnets is through powder metallurgy route. In this process, pre-alloyed powder such as iron-aluminium or cobalt-aluminium may be taken with elemental iron and nickel powders and after mixing in the correct proportion, they are pressed to shape and sintered in vaccume or purely  $\rm H_2$  (24).

An alternative technique involves the milling of alloyed cast material to powder and compacting to solid form by the inclusion of 30 - 40% by volume thermosetting resin powder and then sintered to form magnets (25).

The other possible variation as the formation of alloy from the pure oxides of components through co-reduction process (26). In this case, the alloy is obtained in fine powder form and hence comminution

steps are reduced. Industrial production method follows classification and blending of powders to get a narrow size distribution of fine powders. The powders are made into compacts of required size under magnetic field so that all the fine particles orient their easy direction of magnetization along the applied field direction (26). Then the important stage is sintering of these green bodies to increase their density. It could be solid state sintering or liquid phase sintering.

In liquid phase sintering, a low melting point powders is added and sintered. The low melting point powder melts and fills the gaps and alloys with the high melting point powder particles of the compact to shift the stoichiometry of the final alloys to the right value. By this method, higher density sintered magnets can be made (27). A post sintering heat treatment is usually given to the sintered magnets to improve their magnetic properties.

Another modern technique (28) used for fabrication of Alnico magnet alloys with fully columnar grained structure. In this process, by careful control of thermal gradients during solidification, it is possible to grow long columnar grains with <100> axes lying along the direction of growth. The columnar grain growth is achieved by casting the molten alloy

into a chill face mould with the side walls pre-heated. Solidification is predominantly from the chill face, giving required orientation of the columnar grains. The addition of sulphur, tellurium and carbon helps the columnar growth.

#### 1.13 Application of Alnico Magnets:

The development of almico magnets alloys was followed by a great increase in the industrial applications such as in various machines and in electronic devices. Their applications are based on different functions which they can perform.

These are used to convert the mechanical energy into the electrical energy (or vice-versa) in the magnetic field. This function of magnets is used in small electric motors, dynamos, loudspeakers, microphones, speedometers, magnetos etc. The alnico magnets exert a force on a ferro magnetically soft body. This function of magnets is used in relays, couplings, bearings, clutches, magnetic chucks, clamps and separators (extraction of iron impurities, concentration of ores).

These are also subjected to a direction force exerted by a magnetic field. This function of magnets is used in positioning mechanism, compasses, in some ammeters.

The alnico magnets exert a force on moving charge carriers. This function of magnets is used in magnetrons, travelling wave tube, some cathode rays tube, Hall plates, television devices etc.

#### 1.14 Statement of Problem:

The development of permanent magnets, involving a contineous research for materials, having good magnetic properties, has been going on for past three decades. Many problems are to be solved concerning re-magnetization theory in the magnetically hard materials of the Alnico type. The basic composition and micro addition influence of some elements on the material's magnetic properties have been established.

Nevertheless magnetic properties of the Alnico type alloys observed in practice are far from the those theoretically predicted. There is the still unsolved problem of how to reach higher coercive force and magnetic induction values-closer to the theoretical ones - and how to maximize magnetic induction with the highest coercive force existing at the same time.

Although, in India, these Alnico magnets are in use, they are imported from other country. Yet not much effort has been done to fabricate very good permanent magnets of Alnico type. The magnetic properties of these alloys depend on their composition

and processing parameters. Therefore, investigations on the thermal treatment and chemical composition optimization are carried on with the aim of ensuring the existence of conditions for obtaining the better magnetic properties in Alnico type magnet alloys.

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#### CHAPTER - II

#### THERMITE PROCESS

As the aluminothermic reaction has been used for preparing various Alnico magnet alloys in the present investigation, this chapter deals with the thermodynamic kinetics of the thermite process. A brief description of equipments and materials used in practice and applications of thermite process are also presented.

#### 2.1 Thermite Process:

The preparation of metals and alloys by thermic reduction of metal oxides was discovered by a German Chemist, Hans Goldschmidt (1). He defined thermite reaction as "the reduction of metallic compound by one or several reducing agents in such way that when the mixture (metal compound and reducing agent) is ignited at one place, the reaction continues to go on spontaneously with complete oxidation of the reducing agent, a fluid slag being formed, while the reduced metal or alloy is obtained as a compact regulus.

This aluminothermic reaction is a special case of thermite reaction where aluminium is used as

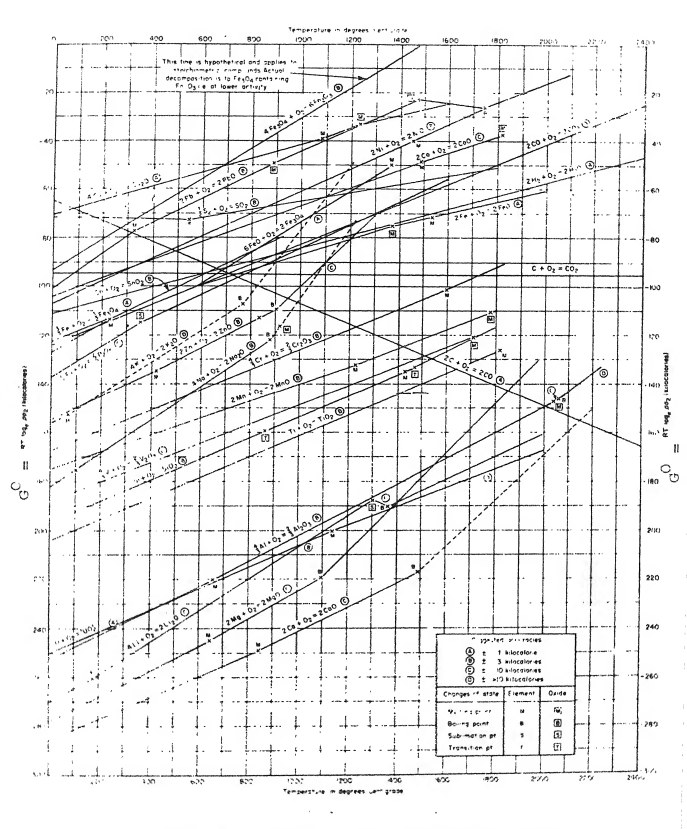


Fig. 2.1: Free energy diagram for oxides

the reducing agent. The use of aluminium is especially attractive due to its availability in high purity at relatively moderate cost, its chemical stability even in finely powdered form and its tendency to reduce most oxides due to having high affinity for oxygen. Fig. (2.1) shows the standard values of free energy of oxide formation as a function of temperature, related to one mole of oxygen (2).

The least stable of the oxides as shown in Fig. (2.1) is that of Mo and the most stable is calcium oxide (CaO). The stability of oxides decreases with increasing temperature with the exception of carbon monoxide (Co). The data shown in Fig. (2.1) can be used to estimate the possibility of using the element as a reductant.

An element, for example Al, which forms a more stable oxide under appropriate conditions can be used to reduce all the metallic oxides shown in Fig. (2.1) expect those of Ca and Zr. The presence of iron or iron oxides can facilitate some reduction processes. Iron dissolves a reduced element, forms a compound with it and thus lowers the activity of that element and removes it from the reaction zone. The general equation for this process is

Me'O + Me" -> Me'O + Me' + Heat

TABLE NO. 2.2

The temperature function of heat capacity (Cp(T) = a+b. $10^{-3}$  T + C. $10^{+5}$  T<sup>-2</sup>), standard value of enthalpy, entropy, heat of fusion, heat of vaporization of metals

Metal	Entropy \(\mathcal{L}\) \$293\(^{\text{OK}}\)	M	<b>д</b>	Heat of fusion	Heat of vaporiza-	H <sub>T</sub> -H <sub>298</sub> OK	Cp=a+] ca:	Cp=a+bx+0 <sup>-3</sup> .T+Cx10 <sup>5</sup> .T- cal/deg/mole	1x105.T
	cal/deg/mo	1e	Value de la companya	(Hf) Kcal/mole			В	q	Ü
A1	6.77	660°3	2467	2.55	70.7	17.93	4.94	2.96	5
A1(1)							7.0	ı	1
ተ ብ	6.49	1539°C	2877°C	3.70	72.00	33.24	4.18	5.92	ı
Fe(1)	•						10.0	ī	ı
Ņ	7.12	1453°C	2902°C	09*9	9° 68	25.26	6.03	-2.49	ì
$N_{1}(1)$							9.02	i	i
Cr	5.68	1898	2665	5.00	81.28	16.93	5.84	2,36	- 88
cr(1)							9.40	ı	ì
8	7.34	1492°C	2677°C	3,64	91.4	20,96	5.26	2.54	ŧ
Co(1)							96.6	ī	ì
<del>S</del>	7.97	1001	2547	3, 12	72.00	13,53	5.41	1.50	1
Cu(1)							7.50	1	

TABLE NO. 2.1

\$tandard free energy of formation of oxides ( G = A + BT) Kcal/mole, Heat of fusion (H<sub>f</sub>) Kcal/mole, Standard enthalpy change ( H<sup>0</sup>293<sup>OK)</sup> Kcal/mole and entropy of oxides (As 293<sup>OK</sup>) cal/deg/mole

at of 3.02930K \$0.298 G=A+BT Kcal/mole slon	<b>C</b> 1	,00 ±00.4 12.24 425.50 ,035	272.7 19.46 230.45 .064	36.7 10.19 43.30 .023	· · · · · · · · · · · · · · · · · · ·	. 57.3 9.03 60.10 .020
Heat of fusion (Hf) Kdal/mole	<b>C</b> 1	52,00 too. too. t	7. 27.2	1 36.7	T Ly.	17.3
4 G <sup>0</sup> 293 <sup>0K</sup> Xcal/mole (+)	129.43	252.65	164.35	33.66	104.35	106,65
Oxide	بع 0ء م	A1203	Cr203	CuO	• OS	Nio

where Me'O is metal oxides and Me" is reducing agent. The reaction is highly exothermic and large negative free energy changes are associated with this reaction as seen from Tables (2.1 to 2.2) and temperature developed during process is expected to be 2525°C.

## 2.2 Thermodynamic Kinetics:

The general equation for the thermite process is

with the equilibrium constant (K) for the reaction as

$$K = \frac{a_{Me'' O :: a_{Me'}}}{a_{Me' O x}}$$

where <sup>a</sup>Me" 0, <sup>a</sup>Me", <sup>a</sup>Me'0, <sup>a</sup>Me' are the activities of reactants and products. The standard isobaric isothermal potential for the reaction is defined as

$$\Delta z^{O} = \Delta z^{O} \text{ Me}^{\prime\prime} \text{ O } - \Delta z^{O} \text{ Me}^{\prime} \text{O} \text{ and}$$

 $\ln K = \frac{-\Delta_G^0}{RT}$ , where  $\Delta_G^0$  is the standard free energy change for the overall reaction. Generally the value of K in the thermite process is in the range of  $10^{10}$  to  $10^{25}$ . Thus for the above reaction to proceed from the left to right, the condition

$$\Delta z^{O}$$
 Me" O <  $\Delta z^{O}$  Me'O

should be satisfied.

The oxygen potential of oxides (3) is defined as

$$\pi_{O(Me'O)} = \Delta z^{O(Me'O)} - RT \ln \frac{a_{Me'}}{a_{Me'O}}$$
+  $\varepsilon (\Delta z) Me'$ , Me'O

and

$$\pi_{O(Me'' \circ)} = \Delta Z^{O}Me'' \circ - RT \ln \frac{a_{Me'' \circ}}{a_{Me'' \circ}}$$
+  $\epsilon(\Delta Z) Me''$ , Me'' o

where  $\epsilon$  (  $\Delta Z)$  Me , MeO indicates the free-energy change due to phase transformation of metals and oxides.

The effectiveness of the reaction is determined by

$$\pi_{O}(\text{Me'O}) > \pi_{O}(\text{Me''O})$$

$$\Delta z^{O} \text{ Me'O} - \text{RT ln} \quad \frac{a_{Me'}}{a_{Me'O}} + \epsilon \text{ (} \Delta Z) \text{ Me'} \text{ , Me'O}$$

> 
$$\Delta Z^{O}$$
 Me" O - RT In  $\frac{a_{Me}}{a_{Me}}$  +  $\epsilon$ (  $\Delta Z$ ) Me", Me"O

The effectiveness of the reaction can be increased by raising  $\pi_{0}$  (Me'O) which is achieved by

- (1) dissolving the Me' formed in the metal solvent and lowering (<sup>a</sup>Me' < 1).</p>
- (2) maintaining  $^{a}$ Me'O  $\Rightarrow$  1 during the process.

The value of  $\pi_0(Me^*0)$  can be lowered by

- (1) using a pure metallic reducer (<sup>a</sup>Me<sup>a</sup> → 1)
- (2) melting and slaging the oxide (Me" O)
- (3) lowering the activity of oxide Me" O through the addition of flux ( ${}^{a}$ Me" O < 1).

These factors promote greater completion of the process and recovery of reduced elements. The successful preparation of high purity magnetic alloys by the Al reduction depends on the use of an excess amount of Al (about 2% of the required Al).

### 2.3 Thermochemical Aspect:

Once the reaction is complete, a metallic phase and a lighter slag will separated out. The slag should be in molten state to prevent the oxidation of metallic phase from atmospheric oxygen. The possible thermite reaction between Al and oxides of various metals (4) is as given below:

Fe<sub>3</sub>0<sub>4</sub> + 8A1 = 9Fe + 
$$4\Lambda l_2$$
0<sub>3</sub>  
 $\Delta G^O$  (2800<sup>OK</sup>) = - 922 + .115 T Kcal/mole  
 $Cr_2$ 0<sub>3</sub> + 2A1 =  $Al_2$ 0<sub>3</sub> + 2Cr  
 $\Delta G^O$ (2800<sup>OK</sup>) = - 145 + .021 T Kcal/mole  
3NiO + 2A1 =  $Al_2$ 0<sub>3</sub> + 3Ni  
 $\Delta G^O$ (2800<sup>OK</sup>) = - 245.2 + .025 T Kcal/mole

$$3^{\circ}$$
0 + 2Al = 3Co + Al<sub>2</sub>0<sub>3</sub>  
 $^{\circ}$ 6 (2800 OK) = - 229.0 + .013 T Kcal/more  
 $^{\circ}$ 8 Cu0 + 2Al = 3Cu + Al<sub>2</sub>0<sub>3</sub>  
 $^{\circ}$ 6 (2800 OK) = = 279.1 + .016 T Kcal/more

when calcium oxide (CuO) is used as flux in the process, then following reaction will occur.

$$3Fe_3O_4 + 8\Lambda 1 + 2CaO = 9Fe + 2(CaO \cdot 2\Lambda 1_2O_3)$$

$$\triangle G^O(2800^{OK} = -922 + .115 \text{ T Kcal/more}$$

$$2Cr_2O_3 + 4\Lambda 1 + CaO = 4Cr + CaO 2\Lambda 1_2O_3$$

$$\triangle G^O(2800^{OK}) = -145 \div .021 \text{ T Kcal/more}$$

$$6NiO + 4\Lambda 1 + CaO = 6Ni + CaO \cdot 2\Lambda 1_2O_3$$

$$\triangle G^O(2800^{OK}) = -245.2 + .025 \text{ T Kcal/more}$$

$$6COO + 4\Lambda 1 + CaO = 6CO + CaO \cdot 2\Lambda 1_2O_3$$

$$\triangle G^O(2800^{OK}) = -229.0 + .013 \text{ T Kcal/mole}$$

$$6CuO + 4\Lambda 1 + CaO = 6Cu + CaO \cdot 2\Lambda 1_2O_3$$

$$\triangle G^O(2800^{OK}) = -229.0 + .013 \text{ T Kcal/mole}$$

$$6CuO + 4\Lambda 1 + CaO = 6Cu + CaO \cdot 2\Lambda 1_2O_3$$

$$\triangle G^O(2800^{OK}) = -279.0 + .016 \text{ T Kcal/mole}.$$

## 2.4 Materials and Equipments:

Materials and equipments required for thermite process can be classified into three headings:

(1) Thermite charge

- (2) Ignitors.
- (3) Thermite crucible and casting mold.

#### 2.4.1 Thermite Charge:

The charge for aluminothermic reduction contains the oxides of metals to be reduced, Al powder alone or in combination with Si. A fluxing agent such as calcium oxide (CaO) or salt-peter (NaNO<sub>3</sub>) is used to decrease the activity of slag thereby increasing the fluidity of slag and thermal blast to either absorb or evalve heat depending upon the process.

Studies conducted on thermite reaction rate by Dubrovinetal (6) proved that the decreasing particle size of oxides results in better mixing and faster rate of reduction. This is due to larger contact area available for reaction.

Al is used in the powdered form and has to be mixed with the oxides. A small excess of Al (2% by weight of total Al) is added to make up for the evaporation losses at high temperatures. For fluxing, lime is mostly used, fluorspar (CaF<sub>2</sub>) and salt peter (NaNO<sub>3</sub>) are also used either in combination with lime or separately. Thermal ballast can be in the form of iron ore, scrap iron and sulphur (7). Sulphur is used when the neat evolved during reaction is not enough to

TABLE NO. 2.3

Ferro-alloy charges All figures refer to weight in kg.

Alloy	Ore	Reducing Agent	Flux	Ballast	Recovery
Fe-Cr	Chrome ore (53 $g$ $cr_2$ 0 $_3$ )2900	A1 350	Saltpetre 30	ı	878
F @•V	Fused $V_2^{O_{\underline{\varsigma}}}$ (75% $V_2^{O_{\underline{\varsigma}}}$ ) 970	Fe-Si 380	Lime 1500	Iron trimming 210	<b>%</b> 866
Fe-T1	ilemite $\operatorname{Conc}_{\bullet}$ (39% Tio <sub>2</sub> ) 1030	Al 400 Fe-Si 10	Lime 100	Scrap 85	70-7 5g
Fe-Nb	Nb ore conc. $(Nb_2O_{\xi})$ 100	Al 32-34	ı	Iron ore 16-17	896
Fe-Zr	Zirconium conc. 100	7167	Lime 30	Iron ore 15	ı
Fe-Mo	Mo ore conc. (518 Mo) 1190	Al-56 Fe-Si 362	Lime 265 Fluorspar 3	Iron ore 290 Steel scrap 210	98.78

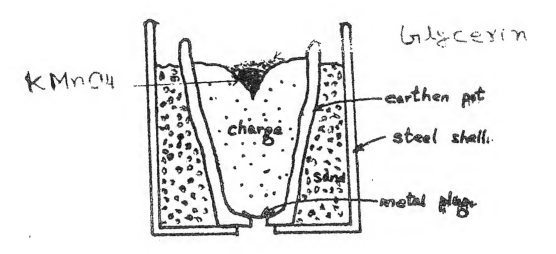
melt the products. Scrap iron or iron ore may added for alloying or to remove excess heat to prevent increase in temperature above the desired level. Table (2.3) summarizes the details of the manufacturing processes for some ferro-alloys.

#### 2.4.2 Ignitors:

The thermite charge requires an ignitor to initiate the reaction. The strip of magnesium is used generally which ignites by applying heat. This method is dangerous as the operator does not get much time to move away from the crucible. Bozzelli (8) had developed a safer ignition system using potassium permanganate crystals and glycerin. The reaction between KMnO<sub>4</sub> and glycerin is an exothermic oxidation and provides enough heat to initiate the thermite reaction. A charge of about 1 kg requires 15 gms KMnO<sub>4</sub> and 5-10c.c. of glycerin. The KMnO<sub>4</sub> is placed in a depression on the surface of hot (200°C) thermite charge and glycerin is then added. The time taken for thermite reaction to initiate is about 30-60 seconds as giving ample time for experimenter to move away from the site of the reaction.

## 2.4.3 Thermite Crucible and Casting Mold:

A cylindrical steel vessel (15 cm dia, 15 cm height), open at the top and a hole of about 2 cm. dia.



Arrangement for firing

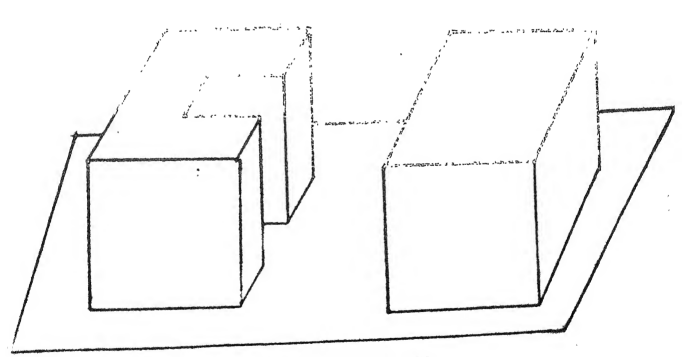


Fig. 2.2 : Exploded view of the Hold

at the bottom centre is used. Holds were provided for lifting the heated vessel. An earthen pot with a hole (.5 to .8 cm dia) drilled through its base and a Al foil is placed over this hole, was used to hold the charge. This was placed inside the vessel and the intervening space was packed with sand. The two holes had been aligned to be one over the other.

Since a large number of ingots had to be made, it was thought that making a sand mold every time would be tedious. Therefore, a permanent mold of cast iron as shown in Fig. (2.2) was used. The stacking of the liquid metal, alloy and slag to the surface of the mold was avoided by coating the walls with graphite.

## 2.5 Applications:

Simple thermochemical concept of thermite process has received much attention in industrial applications due to the following advantages (9).

- (1) The expensive step of preparation of metals or alloys is eliminated.
- (2) Inert gas atmosphere, which is essential if highly reactive metals is handled, may not be required for this process.
- (3) Alloying with a metal whose density is far too low compared to that of base metal might lead to segregation. However, with this process,

this problem is significantly reduced.

A few of the important applications are being discussed in the following paragraphs.

#### 2.5.1 The Preparation of Ferro-Alloys:

The thermite process using alluminothermic reduction is used in the production of ferro-chrome, ferro-vanadium, ferro-titanium, ferro-niobium, ferro-molybdenum and other ferro-alloys (10,11). This insures the production of carbon free products (12).

## 2.5.2 The Preparation of Nuclear Materials:

Aluminium - plutonium alloys which find application as fuel materials in research reactor and as spike elements in power reactors have been largely prepared by aluminothermic reduction (13). The chemical reaction involved in the process is as given below:

PuF<sub>3</sub> + Al = Pu + AlF<sub>3</sub>  

$$\triangle G^{O} 1500^{OK} = -15 \text{ Kcal/mole}$$
  
 $3PuO_{2} + 4Al = 3Pu + 2Al_{2}O_{3}$   
 $\triangle G^{O} 1500^{OK} = -42 \text{ Kcal/mole}$ 

Plutonium yields average 99.8 and product quality is quite high. Similarly alloys of gallium, germanium and silicon can also be prepared by aluminothermic reduction (14).

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## 2.5.2 The Preparation of Nuclear Materials:

Aluminium - plutonium alloys which find application as fuel materials in research reactor and as spike elements in power reactors have been largely prepared by aluminothermic reduction (13). The chemical reaction involved in the process is as given below:

PuF<sub>3</sub> + Al = Pu + AlF<sub>3</sub>  

$$\triangle G^{O} 1500^{OK} = -15 \text{ Kcal/mole}$$
  
 $3PuO_{2} + 4Al = 3Pu + 2Al_{2}O_{3}$   
 $\triangle G^{O} 1500^{OK} = -42 \text{ Kcal/mole}$ 

Plutonium yields average 99.8 and product quality is quite high. Similarly alloys of gallium, germanium and silicon can also be prepared by aluminothermic reduction (14).

## 2.5.3 Aluminothermic Welding:

The reaction between  $\text{Fe}_2^{\text{O}}_3$  and Al is used to weld steel parts.

$$^{\text{Fe}}2^{\text{O}}3$$
 +  $^{2\text{Al}}$  =  $^{2\text{Fe}}$  +  $^{\text{Al}}2^{\text{O}}3$   
 $^{\text{H}}298^{\text{OK}}$  =  $^{2\text{O4}}$  Kcal.

The heat liberated by this reaction is sufficient to obtain liquid iron and slag at temperatures from 1600°C to 1800°C. Thus welding of steel parts is easily possible (15,16).

#### 2.5.4 Production of Refractory Metals:

There have been a number of papers pertaining to the preparation of refractory metals by aluminothermic reduction. Studies on the effect of particle size and relative amount of  $\Lambda l$  powder have been done by Pliner (17). Dubrovin et al (18) studied equilibrium distribution of Cr in the slag phase. Deryabin (19) added chromium oxide to molten aluminum and studied the effects in order to avoid the explosion hazards and lower yields of  $\Lambda l$  -  $Cr_2O_3$  powder reaction.

To prepare high purity vanadium at least 10% Al should be present in V as this suppresses solubility of oxygen. Subsequent removed of volatile Al<sub>2</sub>0 is done by high temperature vaccum sintering. This investigation was carried out by Carlson (20). Carlson and

Schmidt investigated the use of a water cooled crucible as reaction vessel, which resulted in metal of lower  $\rm N_2$  content and consistently higher yields. Wang et al (21) modified this process and adopted it to commercial operation so that Al and oxygen wore removed during consolidation step by electron beam melting. Carlson (20) and Schmidt were able to Co - reduce  $\rm U_2O_5$  containing  $\rm Nb_2O_5$ ,  $\rm Ta_2O_5$ ,  $\rm MoO_3$  or  $\rm TiO_2$  to binary alloys of V.

Wilhelm et al (22) prepared high purity niobium from Nb<sub>2</sub>O<sub>5</sub>. The charge bom is heated to 650°C. Any Al, O<sub>2</sub> or N<sub>2</sub> retained in the reduction product is removed during electron beam melting to give a product containing less than .003% of each impurity. This process is now used commercially.

Because of the low heat reaction of Al with  ${\rm Ta_2}{}^{\rm O}_5$ , Gupta and Jena (7) used sulphur as a thermal booster. The aluminum sulphide also serves as a flux for slag formation by forming an  ${\rm Al_2}{}^{\rm O}_3$  -  ${\rm Al_2}{}^{\rm S}_3$  eutectic. Wilhelm et al (23) accomplished similar results by adding an alloying element such as iron or manganese to form a lower melting eutectic with tantalum. Additional heat is supplied by  ${\rm KclO_4}$  as thermal booster. Iron or manganese with aluminum and oxygen can be removed by electron beam melting.

Mayar and Hubler (24) patented a process for producing niobium or tantalum in an open crucible. The unique feature is the addition of cupric oxide to the charge. Copper which is insoluble with Ta, Nb, wets the metal powder thus forming a protective coating around the individual particles. This adds in the coulesence of powders during subsequent melting or sintering. Copper is then removed by treatment with nitric acid.

Gupta and Jena (25) have been able to achieve yields of upto 70% in the aluminothermic reduction of molybdenum and tungston trioxides. Schmidt (26) has improved the process to increase the yield to 90% at the same time obtaining of 99.98% purity.

## 2.55 Coatings by Aluminothermic Process:

Panteleenko et al (27) used a mixture containing Al<sub>2</sub>O<sub>3</sub> - 38, Al - 18, Cr<sub>2</sub>O<sub>3</sub> - 48 and NHucl - 2% (by weight) for chromizing steels proliminary galvanizing resulted in a 1.3 - 11 fold increase in the chromizing rate and significant increase in wear resistant of the coating. Service life of the galvanized-chromized screws was 3 - 3.7 times longer than uncoated ones. Conventional chromizing without galvanizing gave only 1.6 - 1.8 fold increase in service life.

Lyakhovich et al (23) developed a process for chromizing steel by aluminothermy. A two component (Cr and Si) saturation method was developed for steel using aluminothermic reduction of  $\text{Cr}_2\text{O}_3$  and  $\text{SiO}_2$  in the presence of NHucl activator in  $\text{Al}_2\text{O}_3$  pack. Coatings rich in  $\text{Cr}_3\text{Si}$  and  $\text{Fe}_3\text{Si}$  with micro hardness of 715 - 300 kg/mm² was formed on Armco iron.

Lyakhovich et al (29) investigated thermochemical treatment of austenitic steel. Diffusional saturation of stainless steel with Cr, B, Cr - Si and Cr - Ti was studied using Al reducible  $\text{Cr}_2\text{O}_3$ ,  $\text{B}_2\text{O}_3$ ,  $\text{SiO}_2$  and  $\text{TiO}_2$  as saturation element source optimal cementation treatments increased the wear resistance 20 fold for boronizing, 6-7 for chromosiliconizing, 3 for chromizing and 1.2 times for chromizing with Ti - cementation. Boronizing did not protect the steel from high temperature gas corrosion, but other diffusional coatings increased the oxidation resistance of the steel by 1.2 - 1.5 times.

Yurasko (30) patented a flame spray method for coating metals with a metallothermic mixture containing a thermite component and one or more coating metals.

During spraying the coating metal is superheated to a bonding temperature, resulting in high adherence. The powder blend used was 50% agglomerated NiO with 50% Al as powder of 5µ diameter bonded with sodium silicate.

The mixture was flame sprayed in an oxyacteylene torch on, to a grit blasted steel substract. The resultant coating bond strength of 7245 psi was obtained. For 95% Ni with 5% Al held togather with a phenolic blinder the bond strength was only 5700 psi.

Taguchi (31) has patented coating compositions for molds. Metal oxides and aluminium were coated on to mold to diffuse on cast iron, steel or copper to improve corrosion resistance, heat and abrasion. Thus a coating composition was prepared from 152.2 gm  $\rm Cr_2O_3$ , 53.96 gm Al and a phenolic resin as blinder.

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#### CHAPTER - III

# EXPERIMENTAL PROCEDURES, RESULTS, DISCUSSIONS AND CONCLUSIONS

#### ABSTRACT:

An entirely new method for the preparation of permanent magnetic alloys, using aluminothermic reduction is discussed. The methods of characterizing these alloys, using techniques of optimal metallography, chemical analysis, X-ray diffraction, micro-hardness measurement, magnetization and electrical resisvity measurements are also described. The results of the present investigation are also presented and discussed in this chapter.

## 3.1 EXPERIMENTAL PROCEDURE:

## 3.1.1 Preparation of Alloys:

The materials used in the preparation of alloys were Fe<sub>3</sub>O<sub>4</sub>, obtained locally in the form of flakes with size ranging from (1 to 5 mm), Cr<sub>2</sub>O<sub>3</sub>, NiO, CoO, CuO and Al powder (- 325 mesh), supplied by Sisco Research Laboratory, Bombay. Calcium oxide (CaO) was obtained in the form of lumps, supplied by Galaro Laboratory, Bombay.

Experimental Charges (wt. in gms)
\*\* Refers to Stochiometric.

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(FC	(Fc-Ni-Al-Cr) system								
-		530.44	152.70	283,00	43,86	ŧ	i	272,16	1337.1
2	Fe-208 Ni-58 Al-68 Cr	555, 54	168,00	235,96	52.62	i	ı	270.23	1332.3
m	Fe-248 Ni-58 Al-78 Cr	530, 70	183,24	285,00	61.40	ı	1	269,32	1329.6
ক	Fe-268 Ni-58 Al-38 Cr	505,30	198.54	234.04	70.14	1	ı	263,42	1326.9
Ų,	Fe-278 Ni-58 Al-98 Cr	439.24	206,16	283,92	73.96	2	1	263,14	1326.4
9		472,63	213.78	283.27	87.72	ı	ı	267.70	1325.1
		•	6			7		7	1007
7.	Fe-208 Ni-108 Al-58 Co- 28 Cu	522.36	152.70	237.40	i	33,16	15.00	2/1.59	1631.
ဏ်	Fe-188 Ni-108 Al-108 Co-	489,24	137.46	287.01	ı	76.32	22.56	271.22	1283.6
0	Fe-168 Ni-108 Al-128 30-	430,90	122.16	285,90	1	91.56	30.04	270.13	1280.
10.	Fe-168 Ni-108 Al-158 Co-	458,34	106.92	285.35	í	114.42	37.56	269.66	1272.
11.	Fe-128 Ni-108 A1-208 Co- 68 Cu	431.16	91.62	285.24	1	152.58	45.06	269,55	1275.

followed by addition of about 10 - 15 c.c. glycerin. The reaction starts spontaneously, the melt was cast in the mold.

Most of the characterizations involve regular shape samples of specibic dimension. For this, the specimens of suitable dimensions were cut from the cast ingots with the help of Silicon-carbide cutting wheel and microslicer cutting machine.

### 3.1.2 Heat Treatment

### 3.2.1 Annealing:

The specimens were annealed in  $H_2$  gas with flow rate of 1 c.c./minute at  $1000^{\circ}\text{C} \pm 4^{\circ}\text{C}$  for 3 hours. The specimens were loaded on refractory trays, placed in the middle zone of the furnace. The temperature of the furnace was controlled within  $\pm 5^{\circ}\text{C}$ . The heating rate of the furnace was 3°C/minute and the holding time of the samples at  $1000^{\circ}\text{C}$  was 3 hours. The samples were cooled under hydrogen with rate of  $3^{\circ}\text{C/minute}$ .

### 3.2.2 Tempering:

The annealed specimens were further tempered at 650°C for  $3\frac{1}{2}$  hours in  $H_2$ . The flow rate of  $H_2$  is the same as in the case of annealing.

### 3.2.3 Magnetic Ageing:

The magnetic ageing was carried out in H<sub>2</sub> gas under constant magnetic field using polytronic Electromagnet, Type HEM-75 S.R. No. - 12. For this a specially designed furnace with smaller diameter (in order to fit between the pole pieces of the magnet)

was used. The specimens were aged at 650°C for 1/2 hr. at 1.2 K . Oe under hydrogen stream. At the end specimens were furnace cooled in the applied field in  $H_2$  with flow rate of 1 c.c./minute. The heating rate was 3°C/min and cooling rate was 4°C/minute.

### 3.3 Chemical Analysis:

The large samples of each alloy were cut from the imgots and turned out to remove blow holes and pining defects, developed during casting. The alloy was homogenised at 1000°C for 3 hours in H<sub>2</sub> gas and chemically analysed and the analysed compositions of each alloy were reported in Table (3.2).

## 3.4 Optimal Metallography:

For the metallographic study, the flate surface of the specimens was first polished successively on 1/C through 4/O emery paper taking care that no deep scratches were produced during polishing. The specimens were thoroughly washed in flowing water and again

subjected to polishing on a rotary wheel covered with selvyt cloth using a water slurry of submicron size (0.05µ) alumina as a polishing abrasive. After polishing, the specimens were washed in running water and dried and etched with 10% Nital (a mixture of 10 c.c. of HNO3 and 90 c.c. of methonal). The etching time varied from 2 swabs to 10 swabs depending upon the composition of specimens. The microstructures were examined using Zeiss Universal Metallograph and were recorded on a 35 mm 'Ilford photographic film (160 ISA).

### 3.5 X-ray Diffraction:

The specimens were mounted on perspex sheet of 3 cm square sized with analdite so that flate surface could be ground easily. The polished specimens with flat parallel surface, were studied using a Rich Seigfert 2002 D X-ray Generator and Isodebyeflux diffractometer. The Crka ( $\lambda$  = 2.29f002A°) was used as X-ray source using a graphite monochromater in the path of the diffracted beam. This offered partially background free,  $\beta$ -radiation free, high resolution diffraction patterns for phase(s) analysis, present in the synthesized products.

The diffraction pattern was obtained under following conditions:

Radiation -  $Crk\alpha$  ( $\lambda = 2.291002A^{\circ}$ )

Tube voltage = 40 K.V.

Tube current = 30 mA

Scanning rate =  $3^{\circ}/\min 2^{\circ}$ 

Chart speed = 3 cm/minute

Time constant  $= \cdot 10$ 

Intensity range = 20 K Counts/minute

Angular Range =  $25^{\circ} \le 2.9 < 152^{\circ}$ . of scanning

### 3.6 Microhardness Measurement:

Microhardness of polished samples as mentioned earlier, were measured using Leitz Miniload - 2 Microhardness Tester. The diamond pyramid indentor of the tester was forced onto the surface being tested with a load of 50 P force and the indent was made well within the grain boundary of each phase so that the hardness value obtained did not have any influence due to neighbouring grain. Measuring the length (d) of two diagonals of the indentation with the help of a microscope, the microhardness values were calculated in terms of Vickers Hardness Number using the expression

V.H.N. = 
$$139 \times 10^3 \times \frac{F}{d^2}$$

where F is the applied load defined as F = 50 P = 490.3mN and d is the average length of diagonals (in unit of  $\mu$ m).

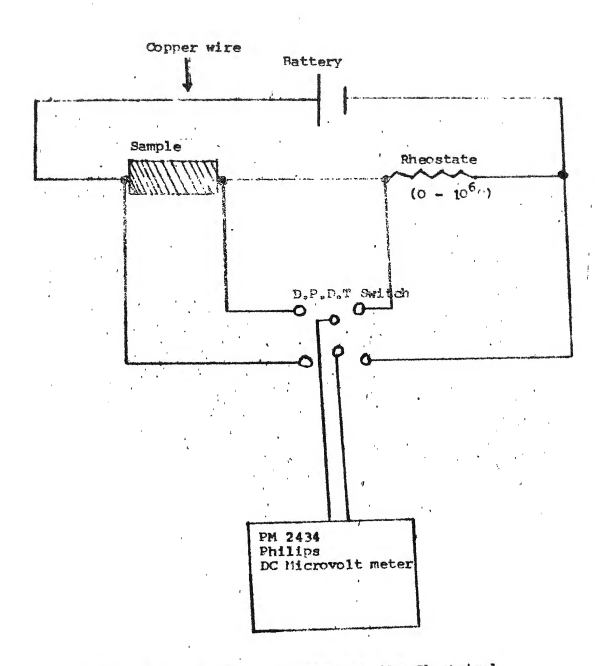


Fig. 3.1: Circuit for measuring the Electrical Resisvity of magnetic materials.

The microhardness of each phase was measured on many grains and the average value was calculated. The microhardness of each alloy was measured in as cast and annealed conditions.

# 3.7 Resisvity Measurement:

The samples were polished. Two copper wire probes were fixed in the opposite faces of polished samples with the help of silver paint and araldite.

The circuit for measuring the resistance of sample is shown in Fig. (3.1). A rheostate was connected in series with the unknown resistance of the sample.

The resistance of the sample was calculated by knowing the total voltage supplied and the voltage drop across the rheostate by PM 2434 Philips D. C. microvoltmeter.

The resisvity of the sample was calculated by calculated resistance of the sample with the following relation

$$R = \int \cdot \frac{1}{A}$$
, where

R is the resistance of sample in  $\mu Q$ ,

f is the resisvity of sample in  $\mu Q$  - cm ,

l is the length of specimen in cm. and A is the contact area of sample in cm<sup>2</sup>.

### 3.8 Magnetic Characterization:

The properties of particular interest for present study of a permanent magnet are saturation magnetization (M<sub>g</sub>), remanence magnetization (M<sub>g</sub>) and intrizsic coercivity (<sup>1</sup>Hc) and dependence of these on various processing parameters.

Samples for magnetic studies were cut into suitable dimensions with the help of microslicer cutting machine provided their maximum dimensions did not exceed 3 cm x 3 cm x 3 cm. APAR 150A Vibrating Sample Magnetometer, supplied by Princeton Applied Research Corp., New Jersey, was used for measurement of M<sub>s</sub>, M<sub>r</sub> and <sup>i</sup>Hc with the maximum attainable field of 10 K. Gauss. The as cast and annealed with tempered with and without magnetic field samples, were studied at room temperature. The principle of the magnetometer is described below in brief.

A sample of the material whose magnetic moment is to be measured is placed in a sample holder. The sample holder is mounted on the end of a rod and suspended and vibrated in the fixed of the electro-magnet. The sample is vibrated in an uniform magnetic field by means of a transformer at a amplitude. The induced field in the sample induces a e.m.f. in a pair of stationary pick up coils. The induced e.m.f. is amplified

by a suitable electromic circuit and converts it into magnetic moment in electro-magnet unit (e.m.u.).

The system is calibrated using the 3/32" dia x 3/32" long cylindrical sample of high purity Ni for which the saturation moment is 55 e.m.u/gm.

The sample is mounted between the electromagnets in a perspex sample holder and is aligned and adjusted there by locating it symmetrically with respect to the detection coils. The main benefit of making this adjustment is that the different samples were adjusted to the same position, relative to the pick up coils, this minimizing the effect of geometry on the result of measurement. The sample is first taken to saturation by increasing the field strength (H) of the electromagnet to its maximum value (10 K . 0e) where the magneto panel meter reads the value of M in e.m.u. The field strength is then reversed to zero and the value of Mr is noted from panel meter. Field direction is then reversed and the field strength is increased slowly till the panel meter reads zero magnetic moment of the sample. Thus we know the value of intrinsic coercivity ( $^{i}$ Hc) at which the magnetic moment of the sample is zero.

#### 3.2 RESULT:

### 3.2.1 Chemical Composition:

The results of the chemical analysis of various alloys (Analysed Composition) is reproduced in Table (3.2) in the last column. The first column represents the composition of starting materials (Nominal Composition) before the thermite ignition. The difference between these two is the result of evaporation of constituent elements or oxides due to high reaction temperature.

### 3.2.2 Optical Metallography:

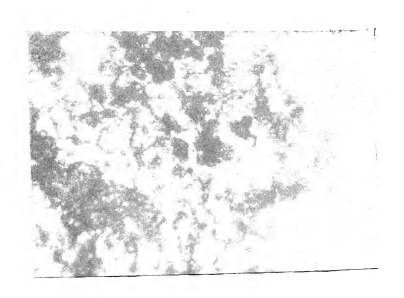
The microstructures of Fe-Ni-Al-Cr system alloys reveal wide-spread precipitates of irregular spaced spherical particles of various shapes and sizes with some inclusions through out the matrix. The grain boundaries were found not to be clearly delineated. The optical micrographs revealed features of phases as one dark black and other grey. The dark areas are due to inhomogeneous distribution of alloying elements in Fematrix. Fig. (3.2) (a, b, c, d, e, f) show microstructures of alloys of this system both as cast and annealed conditions. The annealed microstructures as seen from Fig. (3.2) (d,e,f) show that precipitate particles and inclusions are elongated and highly dispersed to be distinctly resolved.

TABLE 3.2

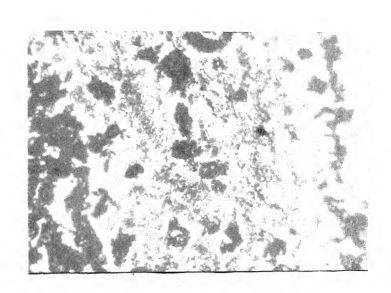
Nominal and inalysed Composition of synthesized alloys

Sample No.	Neminal	inal	COM	posi	tion	composition in wt.pct. Analysed composition in wt. pct.	Analy	rsed com	positi	on in	wt. po	;t.
(Fe-Ni-Al-Cr)	Et I	Fe Fi	7.7	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	8	ch Ch	(Er )	* Ni	12.7	Cr	8 !	Cu
1.	70	2C	44 >	ĸ,	1	1	82,34	11.62	2.54	3.50	ı	ł
2.	67	22	u,	9	ı	1	79.83	12,32	2.63	4.72	ı	ı
°°	64	24	u)	7	1	ı	76,60	14.92	2,52	5,96	ı	1
•	61	56	Κı	ω	ı	ı	74.77	16,12	2.56	6,55	ı	ı
ű.	g)	27	ιΣi	9	1	1	72.32	17.66	2.59	7.43	ı	ı
• 9	57	28	ιζi	10	1	i	68,67	20.22	2.54	8.57	ı	ı
(Fe-Ni-Al-Co-Cu) System												
7.	63	20	¥	ı	ហ	2	77.25	14.02	4.56	i	3.01	1.16
ϡ	63	18	10	1	10	က	74.08	11.95	4.96	ı	6.98	2.03
• 0	23	16	$1^{\circ}$	1	12	₩.	73.17	10.94	4.89	ı	8.04	2,98
10.	. 95	14	10	ı	45	ហ	72.11	8,99	4.89	ı	10.00	4.01
11.	52.	12	10	ı	20	9	70,98	7.10	4.93	ı	12,04	4.95
							-					-

\* Ni content is obtained by difference.

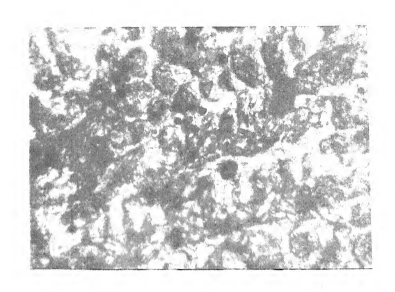


(a) : Microstructure of Fe-11.62% Ni, 2.54% Al. 3.50% Cr as cast (x 300)

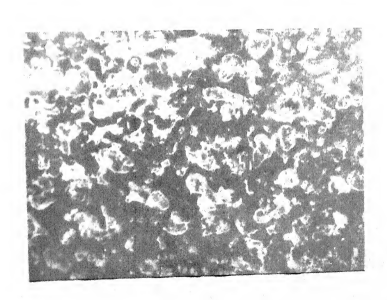


(b) : Microstructure of Fe-11.62% Ni, 2.54% Al, 3.50% Cr, annealed (1000°C, 3hrs) (x 300)

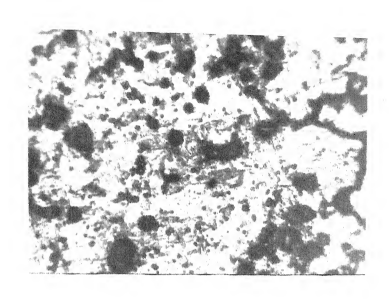
Fig. 3.2: Optical Micrographs of Synthesized Alnico Alloys



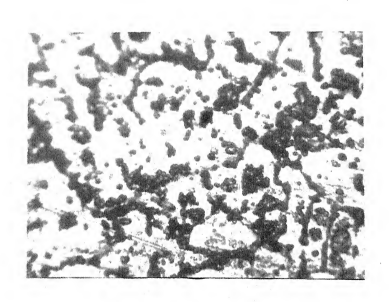
(c) : Microstructure of Fe-16.12% Ni, 2.56% Al, 6.55%Cr as cast (x 300)



(d): Microstructure of Fe-16.12% Ni, 2.56% Al, 6.55% Cr annealed (1000°C, 3 hrs) (x 300)



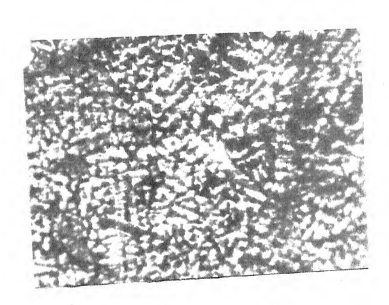
(e): Microstructure of Fe-20.22% Ni, 2.54% Al, 8.57% Cr as cast (x 300)



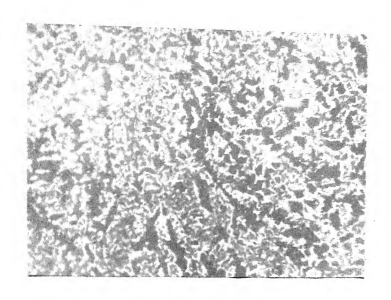
(f): Microstructure of Fe-20.22% Ni, 2.54% Al, 8.57% Cr, annealed (1000°C, 3 hrs) (x 300)



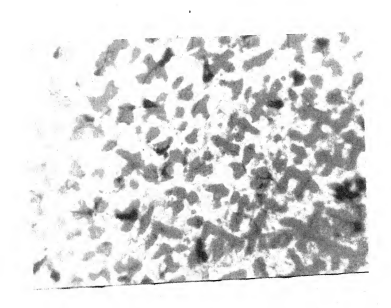
(q): Microstructure of Fe-14.02% Ni, 4.56% Al, 3.01% Co, 1.16% Cu as cast (x 200)



(h): Microstructure of Fe-14.02% Ni, 4.56% Al, 3.01% Co, 1.16% Cu, annealed (1000°C, 3 hrs) (x 200)



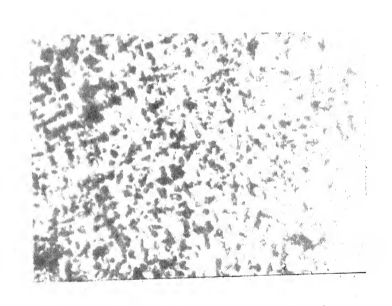
(k) : Hicrostructure of 10.94% Ni, 4.89% Al, 3.04% Co, 2.98% Cu, as cast (x 200)



(1): Microstructure of 10.94% Ni, 4.89% Al, 8.04% Co, 2.98% Cu, annealed (1000°C, 3 hrs) (x 200)



(m): Microstructure of Fe-7.10% Ni, 4.93% Al, 12.04% Co, 4.95% Cu as cast (x 200)



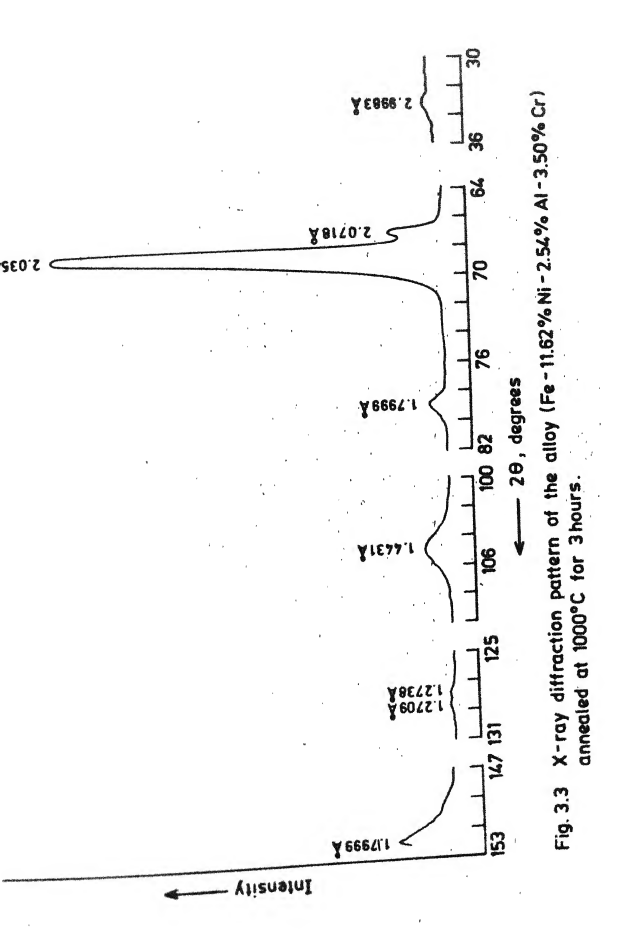
(n): Microstructure of Fe-7.10% Ni, 4.93% Al, 12.04% Co, 4.95% Cu, annealed (1000°C, 3 hrs) (x 200)

The microstructures of Fe-Ni-Al-Co-Cu system alloys show a branch structure of dendrites with some inclusions of unequal size as in cast condition, shown in Fig. (3.2) (g,h,k). These are poorly resolved phase structures. The annealed microstructures of alloys of this system revealed the precipitates of irregularly spaced rod particles of various sizes, oriented with the long axis of the matrix as seen in Fig. (3.2) (1,m,n). The rods tend to group into plate like rows. After annealing, the size of precipitates is uniform, elongated and the precipitates are highly dispersed to be distinctly resolved.

### 3.2.3 X-ray Diffraction:

The results of X-ray diffraction analysis for each of the alloys is presented in Tables (3.3 to 3.13). These tables give the 2  $\theta$ ,  $^{d}$ hkl, relative intensity and the indices of the diffraction lines for the alloys of both systems respectively.

The X-ray diffraction patterns for (Fe-11.62% Ni, 2.54% Al, 3.50% Cr) and (Fe-11.95% Ni, 4.96% Al, 6.93% Co, 2.03% Cu) alloys are shown in Fig. (3.3) and in Fig. (3.4). The reflection pattern observed around the peak is broadened and the peak at 68.5° is indentified as the (220) peak of a b.c.c. phase (Fe<sub>2</sub>Ni Al) with lattice parameter of 5.774A°.



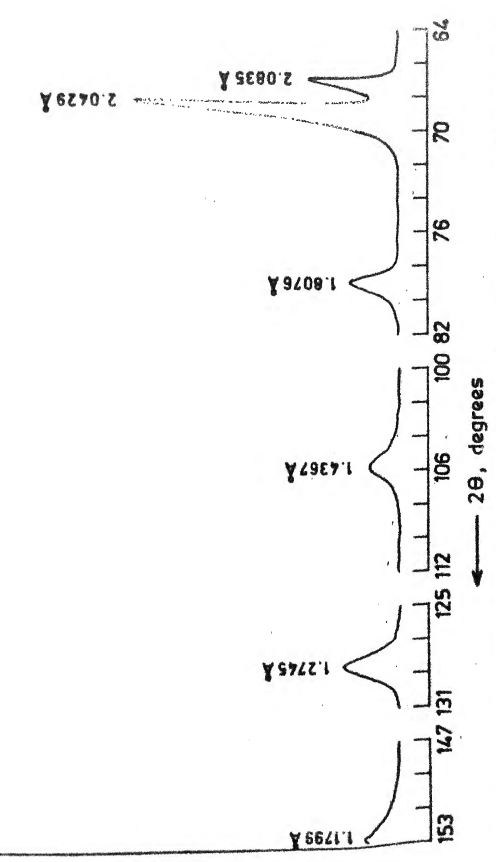


Fig. 3.4 X-ray diffraction pattern of the alloy (Fe-11.95% Ni-4.96% AI-6.98% Co-2.03%Cu) annealed at 1000°C for 3hours.

- YtiznstnI

TABLE 3.3

X-ray Diffraction Pattern of Fe-11.62% Ni, 2.54% Al,

3.50% Cr alloy, Annealed at 1000°C for 3 hrs in H<sub>2</sub>

2 0	d(%°)	I/I <sub>O</sub>	hkl
33.30	3.9979	5	200
66.20	2.0996	18	211
68.50	2.0353	100	220⊭
79.05	1.7999	5	<b>31</b> 0
105.08	1.4431	10	400 <b></b>
123.13	1.2738	4	411,330
129.65	1.2709	4	420
152.25	1.1799	18	332

Crka Radiation  $\lambda = 2.291002 \text{ A}^{\circ}$ ,

To = 10 seconds, C.P.M. = 20 k,

scanning speed = 3°/minute,

chart speed = 3 cm/minute

TABLE 3.4

X-ray diffraction pattern of Fe-12.82% Ni, 2.63% Al,
4.72% Cr alloy, annealed at 1000°C for 3 hrs in H<sub>2</sub>

2 \varTheta	q(v <sub>o</sub> )	I/I <sub>O</sub>	hkl
32.65	4.0753	10	110
58.95	2.3281	<b>2</b> 0	200
62.35	2.2129	10	211
68.46	2.0364	100	220ॠ
79.84	1.7851	20	310
87.08	1.6629	10	222
93.00	1.5925	15	32 1≭
115.10	1.3552	5	400
1/18.50	1.1902	9	411,330
A STATE OF THE PARTY OF THE PAR			

Crk Radiation  $\lambda = 2.29\,1002\,\Lambda^\circ$ , Tc = 10 seconds, C.P.M. = 20 k, Scanning speed = 3 minute in 2 0, chart speed = 3 cm/minute

TABLE 3.5

X-ray diffraction pattern of Fe-14.92% Ni, 2.52% Al,
5.96% Cr alloy, annealed at 1000°C for 3 hrs in H<sub>2</sub>

2 0	₫(¼°)	I/I <sub>O</sub>	hkl
35.50	3.7574	10	110
52.86	2.5736	25	200
59.28	2.3524	30	211¥
63.44	2.0369	100	220⊁
78.95	1.8018	40	310
87.04	1.6635	10	222集
03.12	1.5164	5	321
105.36	1.4404	15	400
125.68	1.2875	20	411, 330
1.27.55	1.1930	20	<b>42</b> 0

Crkm Radiation  $\lambda = 2.291002 \text{ A}^{\circ}$ ,

To = 10 Suconds, C.P.M. = 20 k,

Scanning speed = 3°/min, chart speed = 3 cm/min.

TABLE 3.6

X-ray diffraction pattern of Fe-16.12% Ni, 2.56% Al,
6.55% Cr alloy, annealed at 1000°C for 3 hours in H<sub>2</sub>

2 0	d(V。)	r/r <sub>o</sub>	hkl
34.40	3.3733	10	110
46.36	2.3903	15	200季
56.5	2.4202	19	211
63.40	2.0373	100	220*
75.60	1.3690	25	310
34.04	1.6635	22	222
06.70	1.5330	30	321
105.30	1.4410	10	400 <b>x</b>
123.90	1.2697	19	411, 330
150.30	1.1351	23	. <b>42</b> 0

Crka Rediation  $\lambda = 2.291002 \, \text{A}^{\circ}$ ,

Tc = 10 seconds, C.P.M. = 20 k,

Scanning speed = 3°/min,

Chart speed = 3 cm/min.

TABLE 3.7

X-ray diffraction pattern of Fe-17.66% Ni, 2.59% Al,
7.43% Cr, alloy, annealed at 1000°C for 3 h rs in H<sub>2</sub>

2 0	d(A°)	I/I <sub>O</sub>	hkl
33.8	3.9405	12	110
46.92	2.8832	15	200 <del>1</del>
52.0	2.6131	20	211
68.38	2.0385	100	220⊁
74.7	1.9982	21	310
86.98	1.6644	27	222¥
95.6	1.5463	32	321
105.23	1.4417	25	400
146.2	1.1972	<b>1</b> 9	411, 330

Crkq Radiation  $\lambda = 2.291002 \text{ A}^{\circ}$ 

To = 10 seconds, C.P.M. = 20 k,

Comming speed = 3º/min.

Chart speed = 3 cm/min.

TABLE 3.8

X-ray diffraction pattern of Fe-20.22% Ni, 2.54% Al,
3.57% Or alloy, annealed at 1000°C for 3 hrs in H<sub>2</sub>

3 9	đ (Λ°)	I/I	hkl
2 4. 1.	4.6377	1.4	110
16.79	2.3349	22	200
56.2	2.4320	3	211
63.32	2.0401	100	220 <b>±</b>
75.73	1.8652	16	3 <b>1</b> 0
36.39	1.6631	19	222₩
9.3	1. 5792	20	321
1015.11	1.4425	26	700
134.7	1.2412	15	411,330
117.0	1.1935	12	420

Orka Padiation  $\lambda = 2.291002 \text{ A}^{\circ}$ ,

Chart speed = 3 cm/min,

Scanning speed = 3°/min.

\* Ref : not i and used for lattice parameter calculations.

Fc = 1 comman, C.P.M. = 20 k

TABLE 3.9

X-ray diffraction pattern of Fe-14.02% Ni,

4.56%  $\pm 1$ , 3.01%  $\pm \infty$ , 1.16%  $\pm \infty$  alloy, annealed at 1000°C for 3 hrs in  $\pm 1$ .

2 €	q(v.)	I/I <sub>O</sub>	hkl
62.8	2.1987	32	211
63.43	2.0372	100	220米
73.5	1.9145	16	310
37.05	1.6634	9	222¥
135.	1.2399	17	321
146.3	1.1966	8	400

Crka Radiation  $\lambda = 2.291002 \, \Lambda^{\circ}$ ,

Schming speed = 3°/min, Chart

Speed = 3 cm/min.

\* Refl. 41 no used for lattice parameter calculations

TABLE 3.10

X-ray diffraction pattern of Fe-11.95% Ni, 4.96% Al, 6.93% Co, 2.03% Cu alloy, annealed at 1000°C for 3 hrs. in  $\rm H_2$ 

2 ⊖	g(γ,)	I/I <sub>O</sub>	hkl
66.70	2.0835	34	211
68.2	2.0429	<b>1</b> 00 .	220¥
78.65	1.9076	17	310
105.75	1.4367	9	222
12/3	1.2745	19	32 1×
152.25	1.1799	10	1100

Crka Radiation  $\lambda = 2.291002 \, \Lambda^{\circ}$ ,

To = 10 seconds, C.P.M. = 20 k,

Commin: speed = 3°/min, Chart speed = 3 cm/min.

TABLE 3.11

X-ray diffraction pattern of Fe-10.94% Ni, 4.87 % Al, 8.04% Co, 2.98% Cu, annealed alloy at 1000°C for 3 hrs in H<sub>2</sub>

2 0	q(v <sub>o</sub> )	I/I <sub>O</sub>	hk1
63.5	2.1769	33	211
67.9	2.0506	100	220¥
82.4	1.7391	20	310
96.3	1.6743	25	222¥
97.2	1.5259	· <b>1</b> 0	321
106.6	1.4287	15	400 <b>*</b>

Crka P. Hotton  $\lambda = 2.291002 \text{ A}^{\circ}$ ,

Tc = 10 prevente, C.P.M. = 20 k,

Committe speed = 3°/min,

Chart speed = 3 cm/min

TABLE 3.12

X-ray diffraction pattern of Fe-899% Ni, 4.89% Al, 10% Co. 1.01% Cu annealed alloy at 1000°C ffor 3 hrs. in  $\rm H_2$ 

2 <del>0</del>	d(V₀)	I/I <sub>0</sub>	hkl
36.5	3.6571	27	110
2Fi - 7	2.8890	32	<b>2</b> 00
50.0	2.3264	27	211
68.2	2.0428	100	220¥
77.6	1.8281	21	310
31.3	1.7069	18	222
25.8	1.5112	15	3 <b>2</b> 1 <b>%</b>
107.2	1.4232	9	400
114.5	1.3619	16	411,330¥
125.8	1.2811	9	420

Crka 7. If then  $\lambda = 2.291002 \, \Lambda^{\circ}$ ,

Tc = 10 seconds, C.P.M. = 20 k,

Communitar speed = 3°/min,

Chart speed = 3 cm/min.

\* Ruflant' na used for lattice parameter calculations.

TABLE 3.13

X-ray diffraction pattern of Fe-7.1% Ni, 4.93% Al, 12.04% Co, 4.95% Cu annealed alloy at 1000°C for 3 hrs in  $\rm H_2$ .

		~	
3 H	d(v <sub>e</sub> )	I/I <sub>O</sub>	hkl
3/4.1	3.6976	28	110
26.3	2.9140	31	<b>2</b> 00
59.7	2.3368	29	211
643.1	2.0452	100	220%
77.1	1.8319	23	310
8:.2	1.7087	25	222¥
05.7	1.5451	30	3 <b>2</b> 1#
1.17.00	1.4249	35	400 <b>*</b>
115.24	1.3564	14	411, 330

Crka Radiation - = 2.291002 A°,

Tc = 10 · .... , C.V.M. = 20 k

Scanning speed = 3"/min,

Chart speed = 3 cm/min.

\* Rofl. of the used for lattice parameter calculations.

### 3.2.4 Microhardness Measurement:

The microhardness measurement was carried out to determine the hardness number of the as cast and annualed alloys. The representative microhardness value of each alloy was obtained as an average of several measurement of the second specimen in terms of Vickers Hardness Number.

The result of hardness measurement for all the alloys are shown in Table (3.14). The variation of discrehardness with a specific transition metals (Cr. Co. Cu.) content or total content of precipitating elements is shown in Figs. (3.5 to 3.9). The microhardness of alloys is found to increase linearly with the addition of precipitating elements (single or in combination) in all the cases. Amongst Cu. Co and Cr. the variation in microhardness with the addition of Cu is steepest in comparison to those with Co or Cr. In each case, the annealed alloys have lower V.H.N. than that of the cast alloys.

# 3.3.5 Minctrical Resisvity Measurement:

The measured electrical resisvity of various alloys is presented in Table (3.15). The corresponding resisvity verses alloying element weight percentage is shown in Fig. (3.10) for Fe-Ni-Al-Cr system and in ...

**TAPLE 3.14** 

Microhardness values of Alnico magnet alloys of varying composition as cast and as annealed (at 1000°C for 3 hrs. in  $\rm H_2$ ) conditions.

Sample Wo.	Composition (in wt. pct)	Microhardness* (V.H.N.)	
		cast	Annealed at 1000°C for s in H <sub>2</sub> .
(Fe-Ni-Al-C system	Cr)		
1.	11.62% Ni, 2.54% Al, 3.5% Cr	49 5	435
2.	12.82% Ni, 2.63% Al, 4.72% Cr	540	475
3.	14.92% Ni,2.52% Al, 5.96% Cr	60 5	545
***	16.12%, Ni, 2.56% Al, 6.55% Cr	625	56 5
god Sury on ■	17.66% Ni, 2.59% Al, 7.43% Cr	675	6 <b>2</b> 0
б.	20.22% Ni. 2.54% Al. 8.57% Cr.	725	66 <b>5</b>
(Fe-Ni-Al-C System	(b-(h1)		
~~ <del>*</del>	14.028 Ni, 4.568 Al, 3.018 Co, 1.168 Cu	513	446
3.	11.95% Ni, 4.96% Al, 5.98% Co, 2.03% Cu	625	561
9.	10.94% Ni, 4.87% Al,	660	58 5
10.	8.79% bi, 1.89% Al, 10% Co, 1.01% Cu	<b>72</b> 0	645
11.	7.10% Ni, 1.93% Al, 12.04% Co. 4.95% Cu	780	701

<sup>\*</sup> Without police values indicated above are the average values of his live a measured on several alloys.

TABLE 3.15

Electrical resisvity values of Alnico magnet alloys of varying composition as cast and annealed (at 1000°C for 3 hrs. in H<sub>2</sub>) conditions.

samile Yes.	balance Fe	î Room	Resisvity at Room Temperature (30°C)( $\mu\Omega$ -cm)	
	§ §	l Cast	Annealed at 1000°C for 3 hrs. in H,	
(Fe-Ni-Al-Cr System	)		x 2	
1	11.62% Ni, 2.54% Al, 3.50% Cr.	88	65	
2.	11.82% Ni, 2.63% Al, 4.72% Cr.	91	71	
3.	14.92% Ni, 2.52% Al, 5.96% Cr.	98	75	
₩	16.12% Ni, 2.56% Al, 6.55% Cr	103	<b>7</b> 9	
÷.	17.66% Ni, 2.59% Al. 7.43% Cr.	107	84	
6.	20.22% Ni, 2.54% Al, 3.57% Cr.	110	87	
(Fe-Ni-Al-Co- System	OI)		·	
7.	14.02% Ni, 4.56% Al, 3.01% Co, 1.16% Cu	95	69	
9.	11.95% Ni, 4.96% Al, 6.98% Co, 2.03% Cu	100	74	
9.	10.94% Ni, 4.87% Al, 8.04% Co, 2.98% Ct	103	77	
10.	8.99% Ni, 4.89% Al, 10% Co, 4.01% Cu	104	- <b>7</b> 9	
11.	7.10% Ni, 4.93% Al, 12.04% Co, 4.95% Cu	106	81	

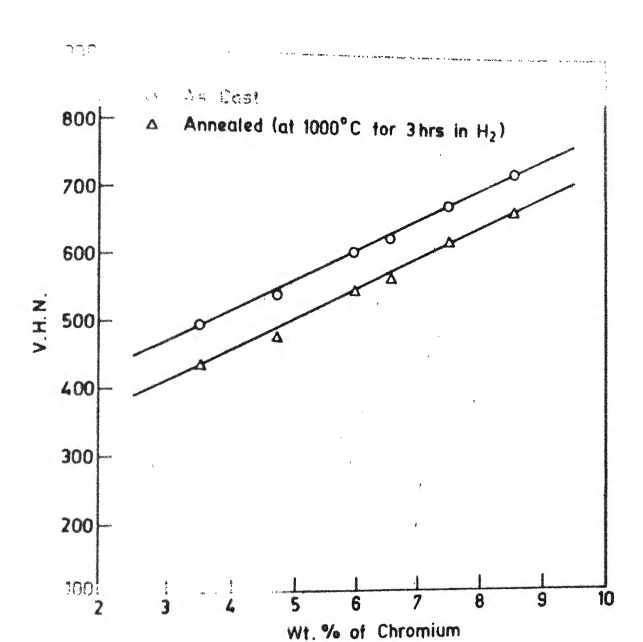


Fig. 35 Variation of V i N. with wt. % of Cr in Fe-Ni-Al-Cr alloys.

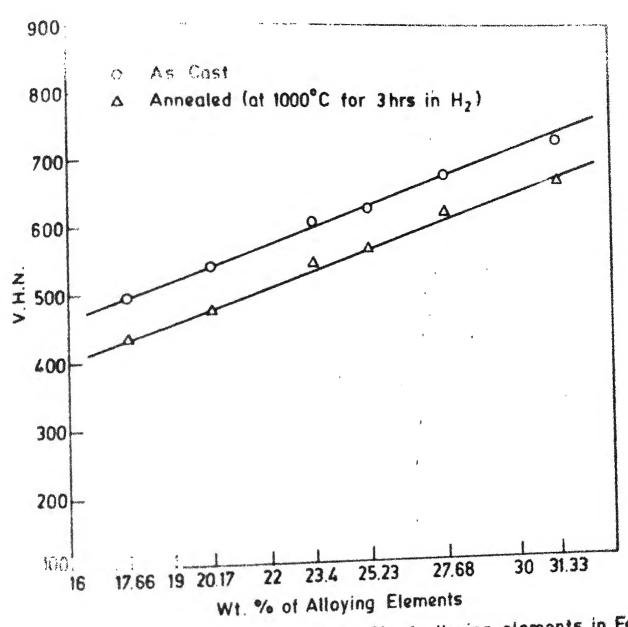


Fig. 3.6 Virialize of V.H.N. with wt. % of alloying elements in Fe-Ni-Al-Cr alloys.

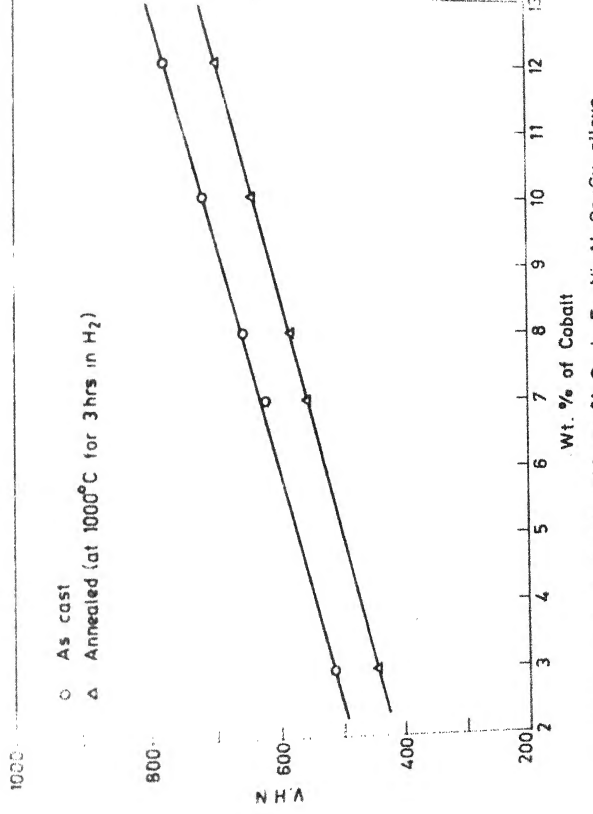


Fig. 3.7 Variation of V.H.N. with wt. % Co in Fe-Ni-AI-Co-Cu alloys.

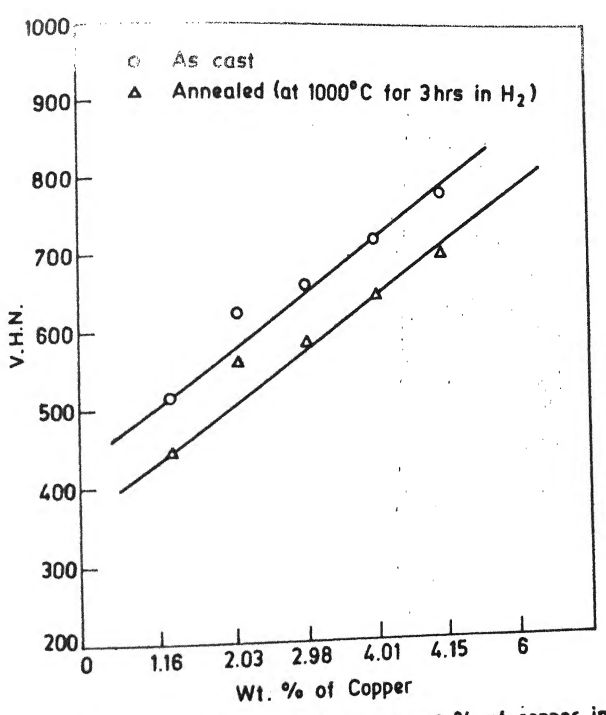


Fig. 3.8 Variation of V.H.N. with wt.% of copper in Fe-Ni-Al-Co-Cu alloys.

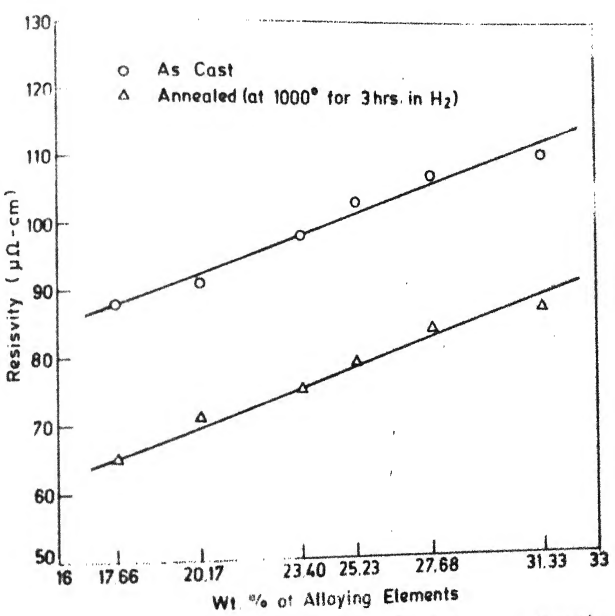
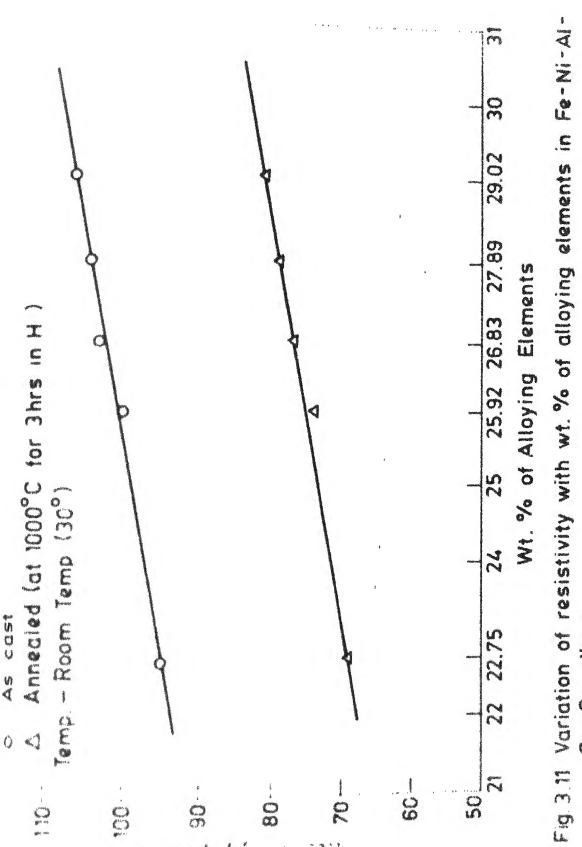


Fig 310 Variation of resisvity with wt. % of alloying elements in Fe-Ni-Al-Cr alloys.



Co-Cu alloys.

Fig. (3.11) for Fe-Mi-Al-Co-Cu system respectively. It is elear that alloying elements inhance the resisvity with increasing addition. In each case, the annealed alloys have lower value of resisvity than that of the cast alloys.

## 3.2.5 Magnetic Measurements:

The aurmary of magnetic measurement for saturation magnetication (Ng), Remanence Magnetization (Mr) and Coercivity (1:1c) is presented in Table (3.16). This is based on experimental study on alloys of (Fe-Ni-Al-Cr) and (Free: i-Al-Co-Cu) systems after giving the following heat tre tments : (1) Annealed at 1000°C for 3 hrs. and tempered at 650°C for 3 hrs. in H2 and (2) Magnetically aged at 6.00°C at 1200 Oe for 1/2 hour in order to determine the dearne to which the magnetic properties can be developed. The magnetimation results are also presented in Figs. (3.12 to 3.13) as plot of M vs H in the first quadrant. The coercive force and remanence magnetization were determined by extrapolation of the demagnetization curve obtained in the second quadrant. Figs. (3.12 to 3.13) show that all the alloys are essentially magnetically saturated in magnetic field ranging from 5.9 to 7.9 KG. For all alloys of both systems, these curves indicate gradual improvement in the changing trend towards loop squareness and increase in saturation magnetization (Mg), remanence magnetization (Mr) and intrinsic coercivity (He) values from the cast state to the magnetically againg state. However, there

5 th

TABLE 3.16

lator	11ed		-						
magnetometer	annealed 1200 /2 hr.	Hc (Ors	red	, ,	9	72	73	79	8 4
	12 0 1	M r e.m.	n/0°0	. 21.81	18.93	17.54	16,84	14.48	13,09
asured	Magnetice at 650°C Crsted fo	M S G · M	1		89 80 82	83.5	795	742	665
em me	for a at h H,	Hc (Ors	200	8	64	69	71	76	₩ ₩
-Ni-Al-Co-Cu) syst	led at 1000°C i and tempered for 35 hrs in	. Mr		. 445	16.94	14. 63	13,56	11.66	10.34
Fe-Ni-	Annealed 3 hrs. al	M.s.e.m.u/		917	855	790	760	700	630
and (		Hc (ors-		52	58	63	99	69	75
(Fe-Ni-Al-Cr)	As cast	м с.с е.ћ.ч/с.с		15,84	13.87	11.92	11.07	و. د تا	7.70
vs of		M S e.m.u/c.c		340	775	720	069	640	575
stic properties of alloys of (Fe-Ni-Al-Cr) and (Fe-Ni-Al-Co-Cu) system measured by	Composition (in wt. pct.) balance Fe.	•		11.62% Ni, 2.54% Al, 3.50% Cr.	12.82% Ni, 2.63% Al, 4.72% Cr	14,92% Ni, 2.52% Ni, 5.96% Cr	16.128 Ni, 2.568 Al, 6.558 Cr	17.66% Ni, 2.59% Al, 7.43% Cr	20.22% Ni, 2.54% Al, 9.57% Cr.
stic pro	O S		i-Al-Cr em	•	.2	m*	- <del>j</del>	ທໍ	ů v

# Contd... Table 3.16

tic proper	tic properties of alloys of (Fe-Ni-Al	-31-Cr)	and (F	e-Mi-7.	1-00-0	u) svst	-Cr) and (Fe-Ni-71-Co-Cu) system measured by magnetometer	by ma	Inetomet	۲
o No	Composition(in wt.pcf.) balance Fe	1	As cast		Annea. 3 hrs. 650°C	led at and for 3	Annealed at 1000°C for 3 hrs. and tempered at 650°C for 3 hrs in H2	Magne annea orste	Magnetically annealed at 650°C orsted for 1/2 hr.	50°C
and the second second second second		e.m. e.	Mr e.m.	14c (ors-	8	Mr 1HC e.m (ors- u/c.c ted	thc (ors-	M S G M C	M M r e.m. e.m.	iHc (ors
l-al-co-cu zetem										
7.	14.02% Ni, 4.56% FI, 3.01% Co, 1.16% Cu	775	18,85	89	980	27.29	78	940	32,90	, <b>8</b>
8	11.95% Ni, 4.96% Al, 6.98% Co, 2.03% Cu.	915	29.05	FT 60	1015	38, 57	86	10.70	46,02	104
•	10.948 Ni, 4.878 Al, 3.048 Co, 2.988 Cu	945	31.96	- <del>1</del> ' 0	1050	44.12	102	1800	53,04	108
10.	8.99% Ni, 4.89% 11, 108 co, 4.018 cu	985	36,51	101	1095	48.18	110	1130	56,35	114
11.	7.10g Ni, 4.93g Al, 12.04g Co, 4.95g Cu.	1020	39.72	105	1125	52,88	113	1130	61, 36	118
							-			

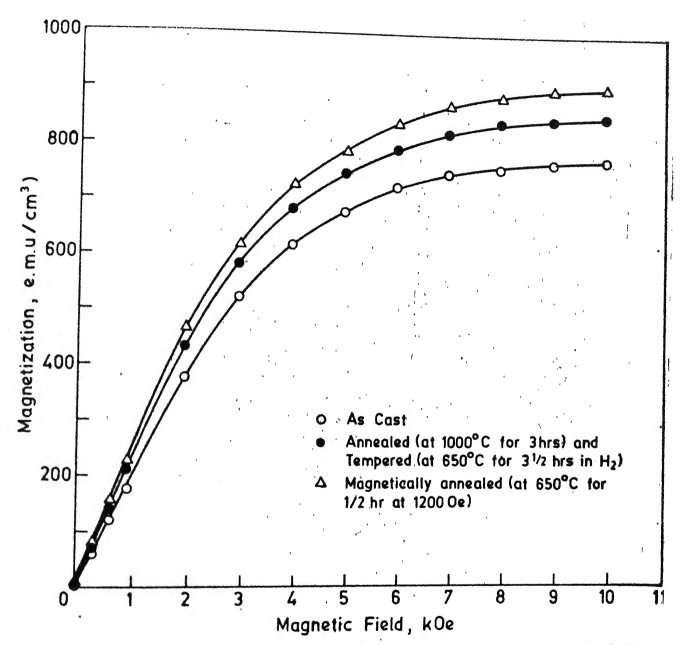


Fig. 3.12 Magnetization curves (First Quadrant) for Fe-14.92% Ni, 2.52% 5.9% Cr

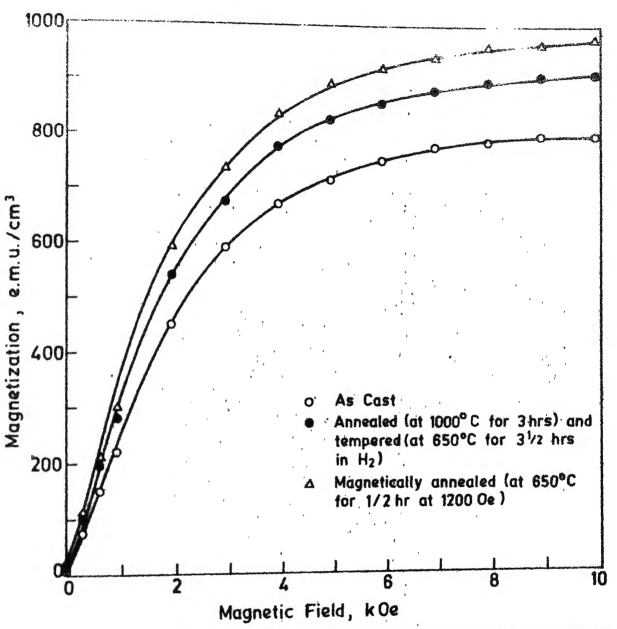


Fig. 3.13 Magnetization curves (First Quadrant) for Fe-14.02% Ni, 4.56% AI, 3.01% Co, 1.16% Cu

is not much difference in the initial magnetization parts drawn for different heat treatment conditions. The gap between magnetization and demagnetization curves for Fe-Ni-Al-Co-Cu system alloys is larger than that of Fe-Ni-Al-Cr system alloys. Thus the alloys of Fe-Ni-Al-Co-Cu system show better approach for loop squareness.

#### 3.3 DISCUSSIONS:

The result obtained in the present investigation are being discussed below under following headings:

- 1. Chemical Composition
- 2. Metallographic Features
- 3. X-ray Diffraction Studies
- 4. Microhardness Measurement
- 5. Resisvity Measurement
- 6. Magnetic Behaviour.

#### 3.3.1 Chemical Composition:

The results of chemical analysis have been presented in Table (3.2). The nominal and analysed composition of all alloys are different. This is attributed to the vaporization of the elemental components, because of very high temperature involved in the process of fabrication. In fact the control of composition of alloys is a serious problem in the present process.

## 3.3.? Metallographic Features:

The microstructures of alloys of (Fe-Ni+Al-Cr) and (Fe-Ni-Al-Co-Cu) systems differ from each-other. Also the structural inhomogenety seen in micrographs of the same type of alloys results due to inhomogenety in temperature distribution between mold wall and the bulk of the melt.

The Fe-Ni-Al-Cr type alloys reveal wide precipitates of irregular shape and size with some inclusions through out the matrix both as in cast and annealed conditions, as shown in Fig. (3.2) (a,b,c,d,e,f,).

During soldification, the sold particles are contineously forming which pass through such zones where the shear forces are weak and the soldified particles may remain in contact with each-other for a some duration without disruption by the fluid forces. This contact duration may be sufficient enough for the formation of much larger particles in size as result of particle coalescence phenomena. The increased content of Cr in the alloy is responsible for increased proportion of inclusions which may be due to insoluble phases such as exides and other intermetallics. Also the high thermal gradient and fast cooling leads to formation of irregular precipitates in size and shape. The microstructures of

Fe-Ni-Al-Co-Cu system alloys show the precipitates in the form of dendritic structure and as irregularly spaced rod particles of various sizes both as in cast and as annealed conditions respectively as shown in Fig. (3.2) (g,h,k,l,m,n). The presence of such particles may be atributed to the effect of copper content on the growth kinetics of  $\alpha'$  phase. Chien and Kattamis (4) have shown that the growth rate of  $\alpha'$  phase in the alloys increases with increase in copper content. The effect of processing conditions on particle size and its distribution has been found to be different for different alloy composition in this system also.

The quality control of alloys of both the systems by the control of microstructures is very complex due to inherent difficulties involved with the thermite process.

## 3.3.3 X-Ray Diffraction Studies:

x-ray diffraction results of the phases present in the alloys of (Fe-Ni-Al-Cr) and (Fe-Ni-Al-Co-Cu) systems are difficult to analysis due to non-equilibrium precipitation of phases during rapid soldification.

There is co-existence of several phases in each system.

Usually for similar system, annealing around 1000°C for several days is recommended in order to obtain a single

very strong line corresponding to a B.C.C. α' (Fe<sub>2</sub> Ni Al) phase with the lattice parameter (a) = 5.774A°. The most intense peak is broad and is observed in the 2 9 range of 68° - 69° (Cr radiation) corresponding to (2.20) reflection. The intensity of this peak did not change visibly with the change in composition of alloys of both systems. Also no systematic change in intensity of other lines with change in composition of the alloys was noticed. Though several second phase precipitates are expected to occur, the X-ray diffraction has revealed only one σ' phase (Fe<sub>2</sub> Ni Al). This α' phase is different in the weakly magnetic matrix.

#### 3.3.4 Microhardness Measurement:

The result of the microhardness measurement of Fe-Ni-Al-Cr and Fe-Ni-Al-Co-Cu systems for as cast and as annealed conditions is presented in Table (3.14).

The data indicates that the addition of Ni, Al, Cr, Co and Cu to Fe - matrix increases the Vickers Hardness Number as shown in Figs. (3.5 to 3.9). These indicate linear dependence of V.H.N. on the concentration of allowing elements in both systems. This can be atributed to distortion introduced in the Fe (B.C.C.) lattice as result of the difference in atomic radii of matrix and the alloying elements. The atomic radii of the elements are Fe = 1.24A°, Fal = 1.43A°, Ni = 1.24A°, Cr = 1.25A°,

Too = 1.25A° and Cu = 1.28A°. The strain field associated with the lattice distortion may offer significant resistance to movement of mobile dislocations during hardness measurement. This may result from the solute atoms. Also the presence of fine precipitates will significant obstacle to dislocation motion. As the matrix distortion and amount of fine precipitates are directly proportional to the concentration of alloying elements, the Vickers Hardness varies accordingly.

#### 3.3.5 Resisvity Measurement:

The result of the resisvity measurement for as cast and as annealed alloys of both the system is presented in Table (3.15). The corresponding variation is shown in Figs.(3.10 to 3.11). These again indicate the linear increase irresisvity with increase in addition of alloying elements. This behaviour can be atributed due to increased electron scattering due to local stress field as a result of solute atoms and second phase particles and dislocations in the Fematrix. All samples of alloys, annealed at 1000°C for 3 hrs. in H<sub>2</sub> exhibit lower values of resisvity than those of cast samples due to relief of residual stress, developed during casting.

## 3.3.6 Magnetic Behaviour:

The magnetization behaviour of those alloys is shown in Figs. (3.12 to 3.13) for samples with different preparation history as indicated in the inset. As is expected, the magnetic properties such as  $\rm M_s$  ,  $\rm M_r$  and  $\rm i_{HC}$  are higher in the case of magnetically aged (at 650°C for 1/2 hr at 1200 0e) alloy in comparison to other conditions. The magnetic ageing in a constant magnetic field through close to curie point is supposed to help the alignment of magnetic domains parallel to the amplied field direction and inducing loop squareness and high values of  $\rm M_s$  ,  $\rm M_r$  and  $\rm ^iHc$ .

All the results presented in Table (3.16) show that the saturation condition for all alloy samples occurs at a reasonably large field (5.9 to 7.9 K . Oe). This may arise due to high demagnetizing factors for small specimens. Moreover, the effect of impurities like carbon on magnetic properties is obscure in the present case. The values of intrinsic coercivity (inc) are found to vary with composition and history of alloy preparation.

## 3.3.6.1 Effect of Cr on Magnetic Properties:

The effect of Cr addition on magnetic properties of Fe-Ni-Al system has been shown in Figs. (3.14 to

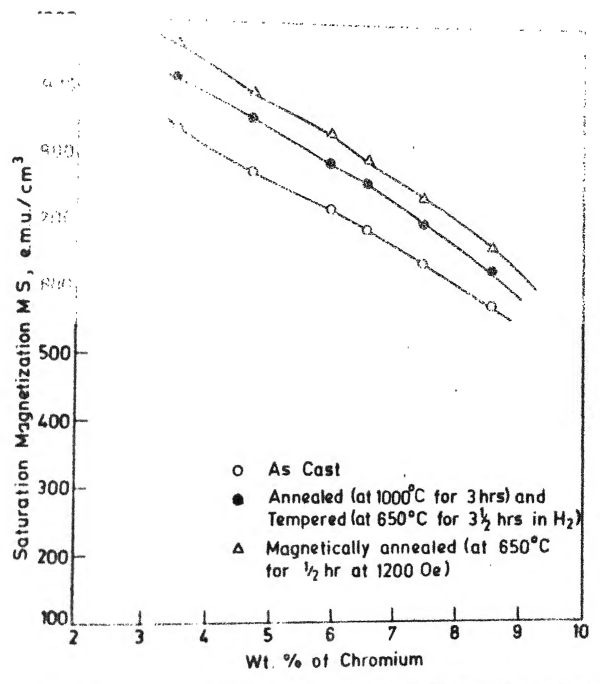


Fig. 3.14 Variation of saturation magnetization MS with weight % of Cr in Fe-Ni-Al-Cr alloys.

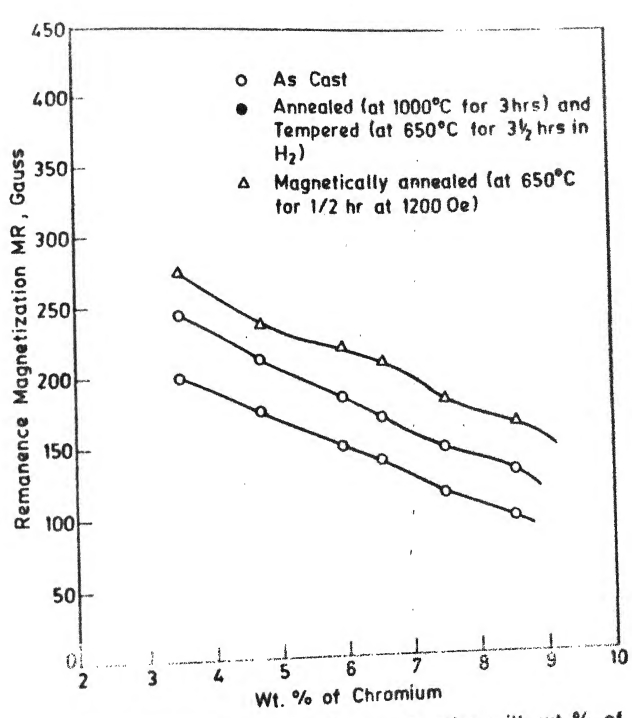


Fig. 3.15 Variation of remanence magnetization with wt. % of Cr in Fe-Ni-Al-Cr alloys.



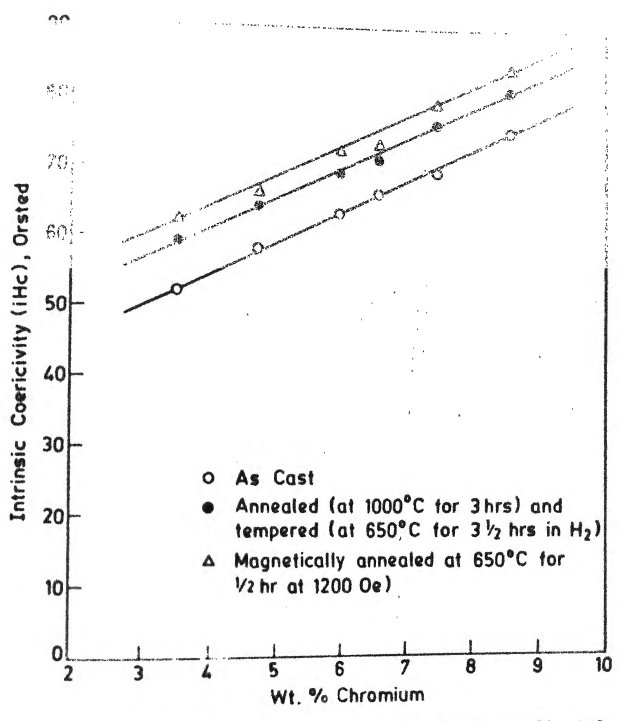


Fig. 3.16 Variation of intrinsic coericivity (iHc) with wt. % of Cr in Fe-Ni-Al-Cr alloys

remanence magnetization  $(M_{\hat{r}})$  are found to decrease with increasing Cr. However, the coercivity is seen to increase with increased Cr content.

A decrease in the values of  $M_s$  and  $M_r$  are explained as following way. A addition of Cr to Fe-Ni-Al system induces the formation of inhomogeneous inclusions with non-unifor distribution with respect to their shape and size. The inclusions causes either the formation of magnetic poles or a stressed domain structure. This reduces the values of  $M_s$  and  $M_r$ . This also increase coercive force by introducing hindrance to magnetization and demagnetization processes due to interaction between domain walls and inclusions (5,6).

The principle of the modern'inclusion' theory was counciated by Neel (7) and some modification in the above theory have been taken up later on in order to account for the effect of non-magnetic inclusions such as Fo<sub>3</sub>C in the ferro-magnetic matrix. Goodengh (3) has discussed the effect of inclusions or precipitates (causing reversal magnetization) on the coercive force in polycrystilline ferro-magnetic materials. His predictions indicate that Hc should increase with 2/3 power of volume of granular inclusions and for lamilar precipitates, Hc should increase linearly with the precipitate volume. The effect of shape and size of

inclusions on iHc has also been considered.

## 3.3.6.2 Effect of Cobalt on Magnetic Properties:

The dependence of magnetic properties of Fe-Ni-Al system on cobalt content is presented through Figs. (3.17 to 3.19). It is evident from Figs. (3.17 to 3.19) that the addition of Co increases M<sub>s</sub>, M<sub>r</sub> and iHc. The increase in M<sub>s</sub> and M<sub>r</sub> values may be atributed to the formation of a ferro-magnetic (Fe-Co) phase in the in the alloys along with a' phase. Though X-ray analysis has not shown definitely the presence of such phase, it is felt that a small amount of ferro-magnetic (Fe-Co) phase is formed which inhances M<sub>s</sub> and M<sub>r</sub> values. The increased intrinsic coercivity may be atributed to the increased shape anisotropy, associated with precipitate particles. Thus the addition of Co has beneficial effect in inhancing magnetic properties of a permanent magnet.

# 3.3.6.3 Effect of Nickel on Magnetic Properties:

The effect of Nickel content on the magnetic properties of Fe-Ni-Al system is shown in Figs. (3.20 to 3.21). The addition of Ni to this system is found to decrease the values of  $M_{\rm S}$  and  $M_{\rm T}$  as shown in Figs. (3.20 to 3.21). This may be atributed to fact that the addition of Ni reduces the average atomic magnetic moment of Ni reduces the average atomic magnetic moment

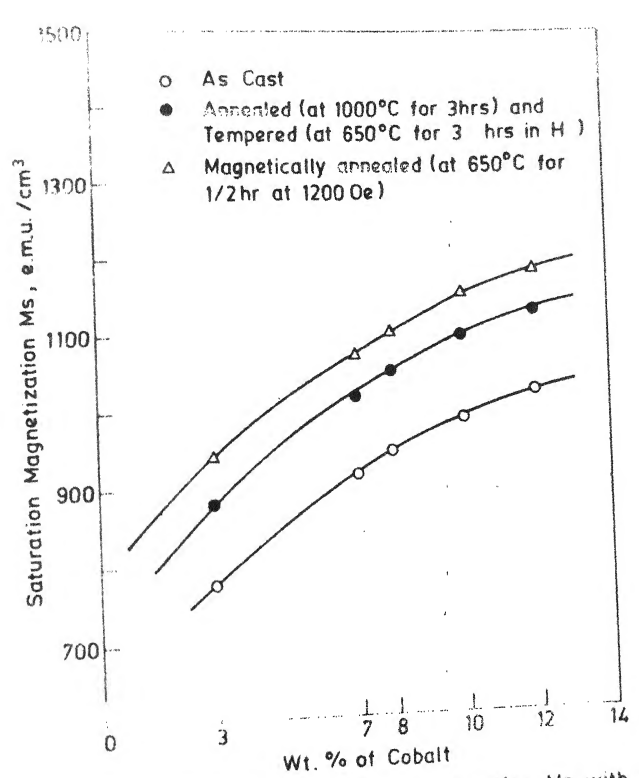


Fig. 3.17 Variation of saturation magnetization Ms with wt.% of Co in Fe-Ni-Al-Co-Cu alloys.

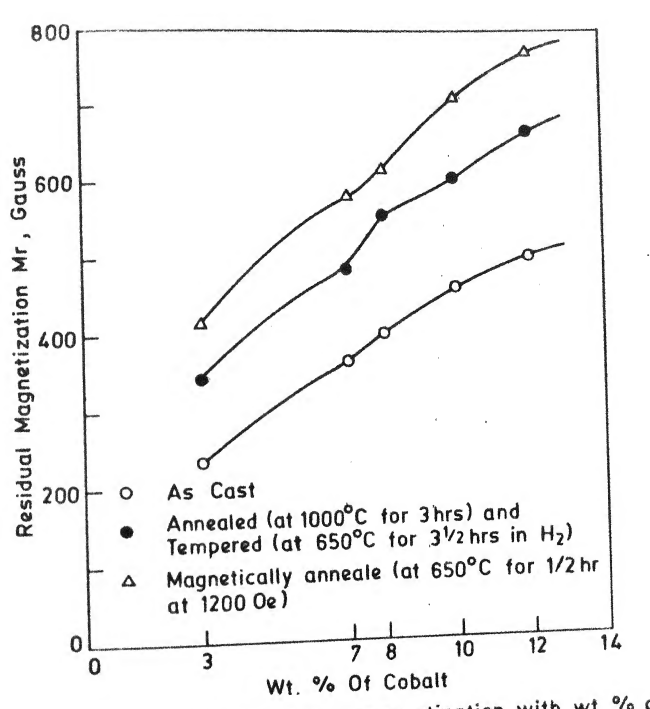


Fig. 3.18 Variation of residual magnetization with wt. % of cobalt in Fe-Ni-Al-Co-Cu alloys.

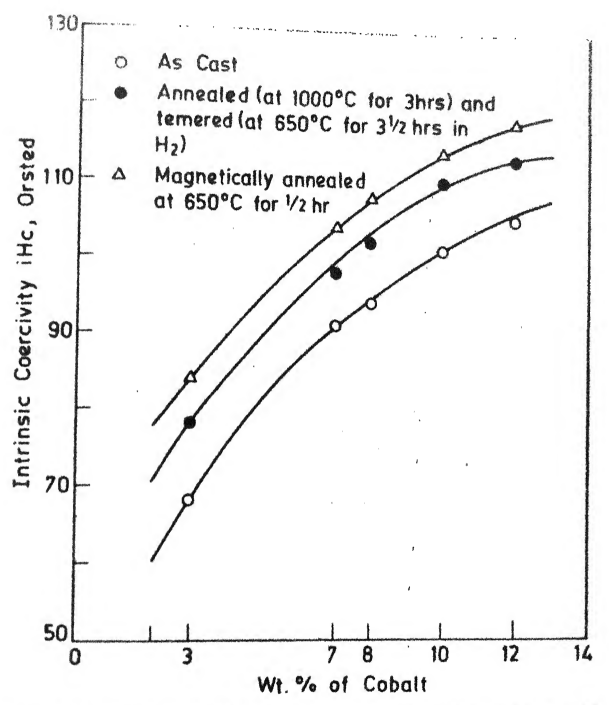


Fig. 3.19 Variation of intrinsic coercivity iHc with wt. % of cobalt in Fe-Ni-Al-Co-Cu alloys.

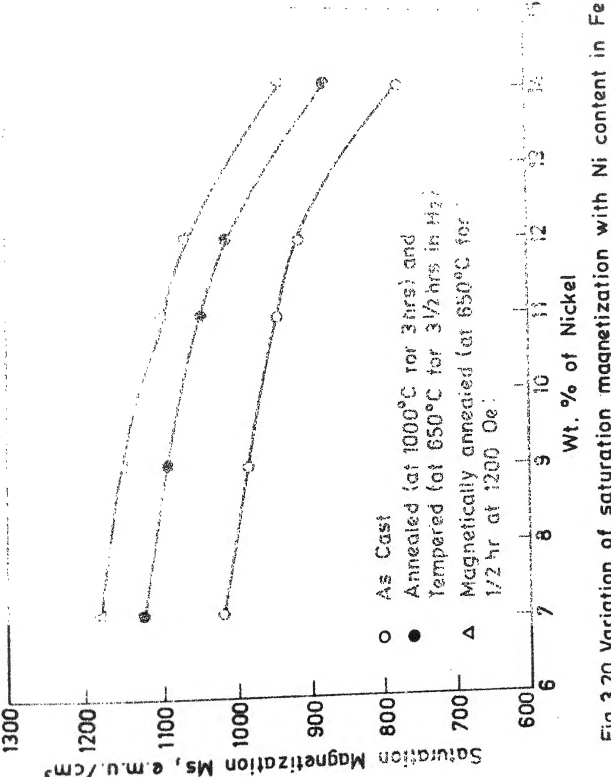


Fig. 3.20 Variation of saturation magnetization with Ni content in Fe-Ni-Al-Co-Cu alloys.

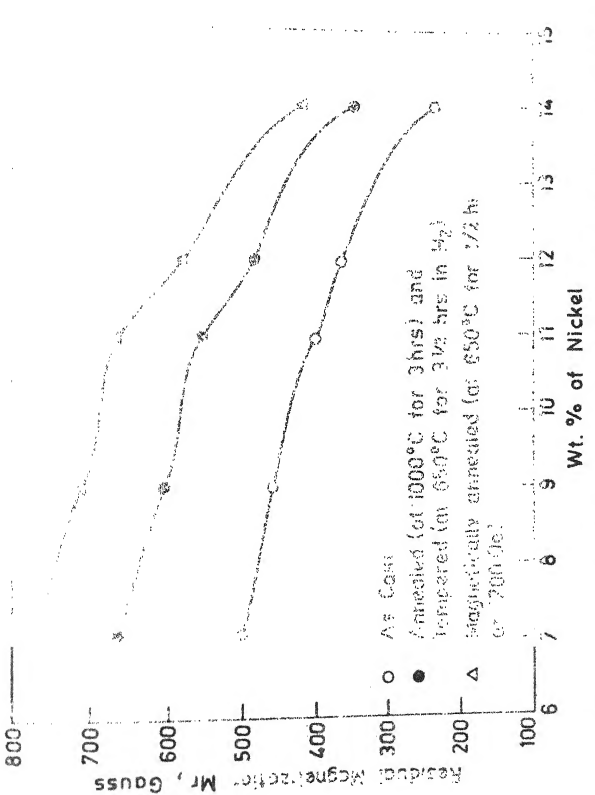


Fig. 3.21 Variation of residual magnetization Mr with Ni content in Fe-Ni-AI-Co-Cu alloys.

is less than that of iron ( $^{m}$ Fe = 2.22  $\mu$ B).

# 3.3.6.4 Effect of Copper on Magnetic Properties:

The effect of Cu content on magnetic properties of Fe-Ni-Al system has been presented in Figs. (3.22 to 3.24). From the above, we find that the addition of Cu increases  $M_{_{\rm S}}$ ,  $M_{_{\rm T}}$  and  $^{\rm i}$ Hc values. The addition of Cu was found to increase the precipitation rate of ferro-magnetic  $\alpha'$  phase in the weak magnetic matrix and induces significant shape anisotropy, giving rise to increased values of  $M_{_{\rm S}}$ ,  $M_{_{\rm T}}$  and  $^{\rm i}$ Hc. The improvement in the values of  $M_{_{\rm S}}$  and  $M_{_{\rm T}}$  due to Cu addition can also be explained using Stoner and Wohlfarth model (9) in which magnetization is assumed to reverse by coherent rotation.

## 3.3.6.5 Effect of Tempering:

The annealed samples of all the alloys were tempered at 650°C for  $3\frac{1}{2}$  hrs. in H $_2$  atmosphere. The results reported in Table (3.16) show that tempering improves M $_s$ , M $_r$  and  $^i$ Hc values in all the cases. Tempering causes greater difference between the spontaneous magnetication of  $\alpha'$  phase and weak magnetic phase due to transfer of Fe atoms from the weak magnetic phase to  $\alpha'$  phase by diffusion process. This results in more effective shape anisotropy of the precipitate particles.



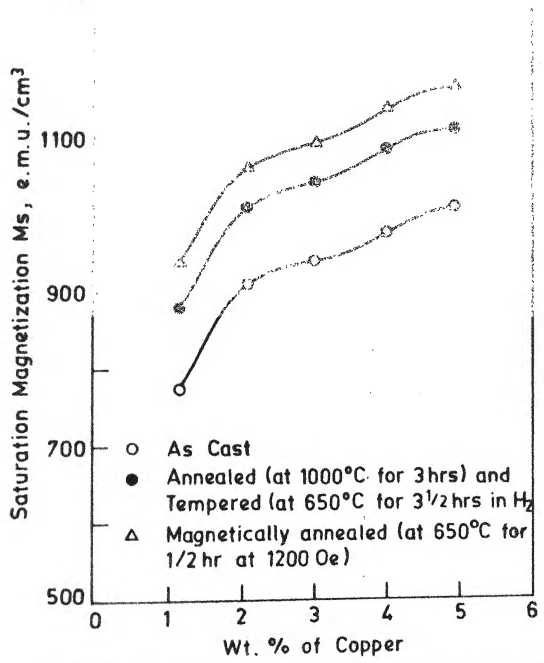


Fig. 3. 22 Variation of saturation magnetization with wt. % of Cu in Fe-Ni-Al-Co-Cu alloys.

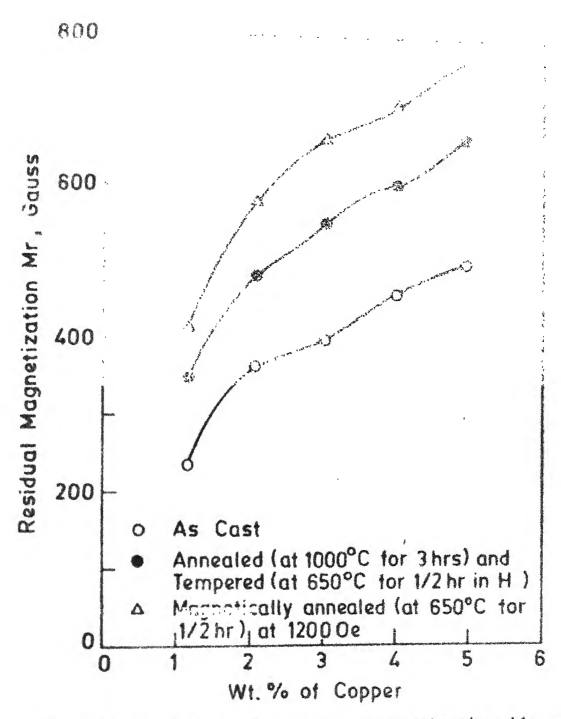


Fig. 3.23 Variation of residual magnetization Mr, with wt. % copper in Fe-Ni-Al-Co-Cu alloys.

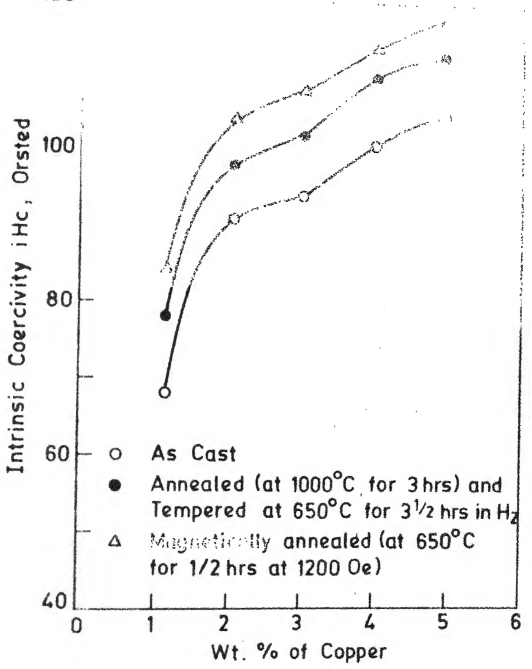


Fig. 3.24 Variation of intrinsic coercivity with wt. % of Cu in Fe-Ni-Al-Co-Cu alloys.

Tempering also causes the clear cut phase separation of  $\alpha'$  precipitates (strong magnetic) and the elongation of  $\alpha'$  particles on the expense of matrix (weak magnetic). This inhances  $^{\rm M}_{\rm S}$ ,  $^{\rm M}_{\rm r}$  and  $^{\rm i}_{\rm H_{\rm C}}$  values.

## 3.3.6.6 Effect of Magnetic Ageing:

The magnetic ageing causes preferential alignment of the spinodally decomposed particles along the field direction or as near as possible, depending on the orientation of [100] of the precipitate particles (10). The elongation of  $\alpha'$  particles inhances shape anisotropy leading to higher value of intrinsic coercivity in the field direction. Thus magnetic ageing results in improved magnetic properties.

## 3.4 CONCLUSIONS:

The present investigation was undertaken to test the conviction of employing metallothermic process for producing hard ferro-magnets using metal oxides in stead of metals. The results of studies on the alloys of both systems led as to the following conclusions:

- It is possibility to produce hard magnets using the thermite process.
- The major difficulty occurs in controlling the composition and microstructure of the alloys.

- The effect of copper addition in Fe-Ni-Al-Co-Cu system is to produce abnormally large rod like precipitates both in the cast and annealed samples.
- The second phase precipitate is a b.c.c. type  $\alpha'$  phase with a = 5.774 $\Lambda^\circ$ . Smaller constituents could not be determined due to their minute amounts.
- 5. Vickers Hardness and electrical resisvity
  were found to increase linearly with addition
  of alloying element concentration.
- The addition of Cr increases the intrinsic coercivity ( $^{i}$ Hc), but lowers  $^{M}$ s and  $^{M}$ r values.
- 7. The addition of Co and Cu improves the values of  $M_s$  ,  $M_r$  and  $^i\mathrm{Hc}$ .
- The annealing (1000°C for 3 hrs.) togather with tempering (650°C for  $3\frac{1}{2}$  hrs in  $H_2$ ) results in higher values of  $M_s$ ,  $M_r$  and  $i_{Hc}$ .
- 9. The magnetic ageing (650°C for 1/2 hr at 1200 Gauss) reveals promising improvement in the values of  $M_s$ ,  $M_r$  and  $i_{Hc}$ .

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